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ERRATUM

In our recent paper on the taxonomy of *Fragilaria* (Kahlert et al. 2019), we relied upon a study of the type material of *F. subconstricta* and *F. tenuistriata* by Tuji and Williams (2008), whose accuracy we had no reason to doubt. Unfortunately, new studies of this material (by Dr Bart Van de Vijver and colleagues, pers. comm., August 2019; see Heudre et al. 2019, published July 2019) indicate that Tuji and Williams inadvertently exchanged these two species in their paper, their account of *F. subconstricta* being based, in fact, on the type material of *F. tenuistriata* and vice versa. Consequently, our paper perpetuates this error and must be corrected on several pages by replacing "*F. subconstricta*" with "*F. tenuistriata*" and vice versa in the abstract, discussion, conclusions, figures, tables, the supplementary figure and its legend, as well as in the supplementary material. Please note that "*F. tenuistriata*" in the results section must not be replaced, as it refers to the clone identification.

In detail, "*Fragilaria subconstricta*" must be replaced by "*F. tenuistriata*" on pages 948 (Abstract), 952 (Fig. 1), 953 (Table 1), 954 (Table 1), 955 (Table 2), 956 (Table 2), 965 (Fig. 7), 966 (4 occurrences; Discussion), 967 (one out of two occurrences; Discussion), 967 (Conclusion), 970 (figure legend to Fig. S3). One sentence needs to be deleted on page 967 (Discussion) because it compares the wrong species with each other and does not make sense when using the correct species descriptions.

In the supplementary material, "*Fragilaria subconstricta*" must be replaced with "*F. tenuistriata*" on page 3 (Table S2), in column "D" (Table S3), Figure S1, Figure S3.

"*Fragilaria tenuistriata*" must be replaced with "*F. subconstricta*" on page 966 (one replacement out of four occurrences; Discussion).

In the supplementary material, "*Fragilaria tenuistriata*" must be replaced with "*F. subconstricta*" on page 3 (Table S2).

The text should have read:

"*Fragilaria tenuistriata*" instead of "*F. subconstricta*" on pages:

952 (Fig. 1)

953 (Table 1)

954 (Table 1)

955 (Table 2)

956 (Table 2)

965 (Fig. 7)

970 (figure legend to Fig. S3)

3 (Table S2)

In column "D" (Table S3)

Figure S1

Figure S3

948 (Abstract)

Our study demonstrated that some species defined on morphological criteria could be confirmed using the *rbcL* chloroplast gene as a genetic marker, for example, *Fragilaria gracilis*, *Fragilaria tenera*, *Fragilaria perminuta*, and *Fragilaria tenuistriata*.

966 & 967 (Discussion)

Tuji and Williams (2008) analyzed the type material of *Fragilaria mesolepta*, *F. tenuistriata*, and *F. subconstricta*, and noted that while *F. mesolepta* has linear valves with a concave fascia and subcapitate ends, both *F. tenuistriata* and *F. subconstricta* have linear valves with a linear fascia and rostrate ends (Table S2). Moreover, *F. subconstricta* has a rimoportula on the sternum of the valve face, whereas *F. tenuistriata* and *F. mesolepta* have their rimoportulae on the mantle-valve face junction. Size, spines (spathulate), and colony formation (long ribbon-like) are similar among all the three taxa. The three strains in the FTNS clade all fit the description of *F. tenuistriata*, whereas the single similar strain 653FraK08 that did not cluster with the other three is *F. mesolepta*, due to the valve outline (constricted median part, subcapitate poles) and the rimoportula on the mantle-valve face junction. Tuji and Williams (2008) noted that *F. tenuistriata* and *F. subconstricta* cannot be separated in LM other than by the position of the rimoportula. In conclusion, this well-supported FTNS clade is clearly *F. tenuistriata* and the single strain not fitting the cluster is *F. mesolepta* because all morphological characters fit these species.

967 (conclusion)

These are: *Fragilaria gracilis* (synonym pro parte: *F. aquaplus*), *F. perminuta*, *F. tenera*, and *F. tenuistriata*.

“*F. subconstricta*” instead of “*F. tenuistriata*” on page:

3 (Table S2)

We apologize for this error.

REFERENCE

Kahlert, M., Kelly, M.G., Mann, D.G., Rimet, F., Sato, S., Bouchez, A. & Keck, F. 2019. Connecting the morphological and molecular species concepts to facilitate species identification within the genus *Fragilaria* (Bacillariophyta). *J. Phycol.* 55:948-70.

1 **CONNECTING THE MORPHOLOGICAL AND MOLECULAR SPECIES CONCEPTS TO FACILITATE**
2 **SPECIES IDENTIFICATION WITHIN THE GENUS *FRAGILARIA* (BACILLARIOPHYTA)¹**

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14 **Running title: UNRAVELLING THE GENUS *FRAGILARIA***

15

16

Abstract

This paper evaluates taxonomic relationships in *Fragilaria* sensu stricto, an abundant and ecologically important diatom genus, taking advantage of cultured and DNA-barcoded material. The ultimate goal is to facilitate identification of European taxa within this complex, providing a unified view on morphological, molecular and ecological relationships. This will simplify both research and environmental assessment, based on either microscopical or molecular analyses. There is general agreement that the separation of species within the group of *Fragilaria* is difficult and consequent confusion can blur potentially important ecological distinctions between species. Our study demonstrated that some species defined on morphological criteria could be confirmed using *rbcL* chloroplast gene as a genetic marker, e.g. *F. gracilis* Østrup, *F. tenera* (W. Sm.) Lange-Bert., *F. perminuta* (Grunow) Lange-Bert. by Tuji & Williams and *F. subconstricta* Østrup. However, even for those species, preliminary identifications based on morphology were often different to those finally established by phylogenetic clustering. A deeper study of morphological characters using both light and scanning electron microscopy confirmed that identification of *Fragilaria* by light microscopy is indeed difficult, and that phylogeny based on DNA barcodes may be a more precise means of differentiating species. Based on molecular and morphological data, we describe three new species: *Fragilaria agnesiae*, *Fragilaria heatherae*, and *Fragilaria joachimii*. Finally, we found well-defined subgroups within one morphological species (*F. gracilis*), whose biogeography and ecology require further study.

Key index words: cryptic taxa, DNA barcoding, *Fragilaria*, integrative taxonomy, *rbcL*, species delimitation, taxonomy

List of abbreviations: RBGE, Royal Botanic Garden Edinburgh, UK. TCC, Thonon Culture Collection, France. IPS, Indice de Polluosensibilité Spécifique. TDI, Trophic Diatom Index.

42 LM, light microscopy. SEM, scanning electron microscopy. HTS, high-throughput
43 sequencing. NCBI, National Center for Biotechnology Information. MOTU, molecular
44 operational taxonomic unit.

45 Working name abbreviations for *Fragilaria* groups, assigned to preliminary identifications:
46 FCAP, *F. capucina*. FGRA, *F. gracilis*. FPEM, *F. perminuta*. FTEN, *F. tenera*. FTNS, *F.*
47 *tenuistriata*. FVAU, *F. vaucheriae*. FCRNAPA, *F. crotonensis*-*F. nanoides*-*F. pararumpens*.

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50 Introduction

51 This paper evaluates taxonomic relationships in *Fragilaria* sensu stricto, an abundant and
52 ecologically important diatom genus, taking advantage of cultured and barcoded material
53 from two different collections (Royal Botanic Garden Edinburgh, UK (RBGE); Thonon
54 Culture Collection, France (TCC)). The goal of this paper is to facilitate identification of
55 European taxa within this complex, providing a unified view on morphological, molecular
56 and ecological relationships and an improved basis for biomonitoring using DNA
57 metabarcoding (cf. Vasselon et al. 2017, Kelly et al. 2018).

58 Species of *Fragilaria* sensu stricto are often an important part of the diatom assemblage in
59 freshwater ecosystems, spanning a range of ecological conditions. For example, diatoms
60 identified as *F. capucina* var. *vaucheriae* (Kütz.) J. B. Petersen were present in 642 of 2170
61 samples collected in the eastern part of France (Rimet and Bouchez 2012a), whilst *Fragilaria*
62 *gracilis* Østrup was the third most commonly recorded diatom taxon in 1013 samples
63 collected in Sweden (Kahlert 2011).

64 It is important to separate different taxa from each other because they can give different
65 ecological information. For example, of the 37 freshwater *Fragilaria* taxa in the Swedish
66 standard checklist of diatoms for biomonitoring (Kahlert et al. 2017), their Indice de
67 Polluosensibilité Spécifique (IPS) sensitivity values (Cemagref 1982) range from 3.4,
68 indicating moderate pollution or eutrophication, to 5, indicating no pollution and very high
69 water quality. The Trophic Diatom Index (TDI) sensitivity values (Kelly and Whitton 1995)
70 for *Fragilaria* taxa range from 1, indicating that a taxon is mainly present at very low
71 phosphorus concentrations to 3, indicating that a taxon can tolerate moderate phosphorus
72 enrichment. The pH preferences of *Fragilaria* taxa range from acidophilous to alkaliphilous
73 (Van Dam et al. 1994). Some *Fragilaria* species form colonies whilst others are found as
74 single cells only (Rimet and Bouchez 2012b, Lange-Bertalot and Ulrich 2014), and whereas

75 many live amongst the benthos of rivers or lakes, others are planktonic (e.g. Rimet and
76 Bouchez 2012b) or maybe even alternate between the two (e.g. Lange-Bertalot and Ulrich
77 2014).

78 However, there is general agreement that separation of species within *Fragilaria* is
79 challenging. Krammer and Lange-Bertalot (1991) stated that the taxa used up to then were
80 often difficult to separate, sometimes due to issues with original type description and often
81 because, even when type descriptions were adequate, morphological characteristics overlap.
82 Since then, *Fragilaria* has received more attention: several groups of species have been split
83 off into other genera, type material of many taxa has been studied, new taxa have been
84 described, and comparisons between taxa have been made (Tuji and Williams 2013, Lange-
85 Bertalot and Ulrich 2014, Delgado et al. 2015, Wetzel and Ector 2015, Almeida et al. 2016).
86 Problems remain, however, despite these studies: there are no or only limited descriptions of
87 the type for several species, scanning electron microscopy (SEM) observations are sometimes
88 missing, and there may be no descriptions of traits such as colony formation, or habitat. It is
89 also not helpful that comparisons of new with existing species often do not take all similar
90 taxa into account, leaving readers wondering if certain taxa are no longer considered to be
91 valid: for instance, *F. nanoides* Lange-Bert., is not mentioned in Lange-Bertalot and Ulrich
92 (2014).

93 Different laboratories have developed their own strategies to handle these issues, leading to
94 species concepts and descriptions that do not necessarily resemble each other. As an example,
95 the common species *F. vaucheriae* (Kütz.) J. B. Petersen – known to be tolerant to high
96 nutrient concentrations, organic pollution and even pesticides in contrast to other species
97 within the *Fragilaria capucina* complex sensu Krammer and Lange-Bertalot (1991), which
98 are seen as more sensitive (Cemagref 1982, Lecoinge et al. 1993, Kelly and Whitton 1995,
99 Larras et al. 2013, Lecoinge 2018) – is differentiated using different criteria by different

authors: The original type material, studied by Wetzel and Ector (2015), shows a taxon with a length of 14–50 μm , a width of 3.8–5.1 μm , and 11–14 striae per 10 μm . However, Hofmann et al. (2011) report 9–14 striae per 10 μm whilst the online Diatoms of the United States (Morales 2010) states 14–16 striae per 10 μm . In a compilation aimed to harmonize identification of diatoms from the Baltic Sea, *F. vaucheriae* is described as having a width of 2.5–4 μm (Snoeijs and Potapova, 1995) and, in the European Diatom Database (Battarbee et al. 2001), *F. vaucheriae* has been merged with *F. rumpens* (Kütz.) G. W. F. Carlson. The situation gets more complicated because Krammer and Lange-Bertalot (1991) omitted *F. pectinalis* (O. F. Müll.) Lyngb. which is narrower and has finer striae than *F. vaucheriae* whilst Wetzel and Ector (2015) have described *F. microvaucheriae* C. E. Wetzel & Ector sp. nov.

Fragilaria gracilis presents another example of confusion within this genus. Even though type material was studied and compared with similar taxa by Tuji (2007) and Lange-Bertalot and Ulrich (2014), its separation from other species has not necessarily become clearer. Lange-Bertalot and Ulrich (2014) give a key where the first character separating *F. gracilis* from other similar taxa is “no formation of colonies”. However, most diatom analyses are based on prepared slides with limited opportunities to check the unprepared sample for colony-formation. Moreover, even if there is access to fresh material, it can be difficult to directly link the unprepared specimens of *Fragilaria* to identifiable cleaned (i.e. oxidized) specimens on prepared slides or SEM stubs if there are several similar taxa present in a single sample (which is often the case). So colony formation, even if helpful for a species description, and important in ecology, is not a very helpful characteristic for routine diatom identification. The next criterion used by Lange-Bertalot and Ulrich (2014) is valve shape (“needle-shaped”), which then leads to a choice between “valve ends narrowly rounded, never subcapitate or capitate”, which would identify the diatom as *F. saxoplanktonica* Lange-Bert.

et Ulrich, or “valve ends capitate or at least weakly subcapitate, sometimes protracted”, which would be other species, among them *F. gracilis*. Tuji (2007), on the other hand, studied the type material of *F. gracilis* and reported that “the form of apex [of *F. gracilis*] can vary from acute to subcapitate”. So again this character is not really helpful in routine analysis. Finally, separation of *F. gracilis* from the recently described species *F. aquaplus* Lange-Bert. et S. Ulrich is based mainly on the number of striae (*F. gracilis* 19.5–21.5 per 10 μm , *F. aquaplus* 22–24 per 10 μm), which is a good character to use for routine analysis because it can be easily measured. However, the additional criterion of the striae “opposite or alternating” (*F. gracilis*) versus “mostly opposite” (*F. aquaplus*) is again weak, as the qualifier “mostly” requires the analyst to make value judgements about whether an exception to a general rule indicates a different species or simply phenotypic variation within a population. In summary, different experts analyzing diatom samples probably have a quite different picture of *F. vaucheriae* or *F. gracilis* and relatives, leading to problems when taxa lists from different laboratories, projects and countries are compared (e.g. Kahlert et al. 2009).

While traditional descriptions of diatom species and identification aids have been based solely on morphological characters, we now have an additional tool to gain deeper insights into taxonomy: DNA sequences. Using DNA sequences has provided a completely new set of characters to compare, helping to unravel species boundaries and names, and giving a level of clarity about species boundaries that was extremely difficult to obtain previously (Amato et al. 2007, Sarno et al. 2007, Evans et al. 2008, Pouličková et al. 2010, Vanormelingen et al. 2013). Appropriate short regions within selected genetic markers can be used as ‘DNA barcodes’ for identification (e.g. Mann et al. 2010), especially during applications of high-throughput sequencing (HTS) technologies for DNA metabarcoding. Analyzing environmental samples with DNA metabarcoding is already being applied in ecological assessment and research (Vasselon et al. 2017, Kelly et al. 2018, Rimet et al. 2018b) and will

also give new insights into diatom taxonomy as researchers investigate the biological meaning and phylogenetic significance of clusters of DNA sequences (molecular operational taxonomic units, MOTUs).

In this study we aim to unravel *Fragilaria* species taxonomy and names by comparing a phylogenetic tree created from molecular data with observed morphological characters, and with a list of morphological features compiled from published species descriptions and type material. Our goal is to ensure consistency in the use of species names, to supply reference sequences (and hence DNA barcodes for HTS analyses), and where possible to identify diagnostic morphological features. We aim to define monophyletic taxa based on molecular data because paraphyletic taxa are almost impossible to detect with DNA metabarcoding. This, in turn, will give more accuracy and robustness to ecological values of *Fragilaria* species, facilitating the development of DNA-based approaches for ecological assessment using diatoms.

Material and methods

Studied strains

The strains studied come from two different culture collections: the Royal Botanic Garden Edinburgh (RBGE) and the Thonon Culture Collection (TCC), France. They were initially identified to species using the light microscope (LM) and common diatom identification literature, before being sequenced and investigated by SEM. For this study, we selected all available strains of both collections identified as belonging to the genus *Fragilaria* and the neighbor genus *Ulnaria*. For detailed information on the studied strains see Table S1, and Rimet et al. (2016), Kelly et al. (2018), Mougin et al. (2018), and Rimet et al. (2018a).

128 strains were available from RBGE, collected and isolated from Scotland and England in 2012 (hereafter United Kingdom, UK); all were sequenced for *rbcL* as described by Kelly et al. (2018). The isolates were cultured only as long as required to generate enough material for DNA extraction. For voucher material, subsamples were dried onto small round cover slips (13 mm diameter for SEM, 18 mm diameter for LM). For LM, the diatoms adhering to the cover-slips were cleaned in situ using hot 70% HNO₃ and then washed with deionized water before mounting in Naphrax (Brunel Microscopes, Chippenham, UK). Light micrographs were taken (by S. Sato and D. Mann, for preliminary identification for the biomonitoring project of Kelly et al. 2018) prior to the present project but no SEM observations of the *Fragilaria* clones had been made. DNA extraction and *rbcL* sequencing of the RBGE clones is described by Kelly et al. (2018).

The TCC collection hosts isolates from different countries and geographical regions. Those used in our study were from Europe (France, Italy, Portugal, Luxembourg, Sweden), and a few from the over-sea territories of France (Ile de La Réunion). The TCC database also includes information about curated strains from NCBI (National Center for Biotechnology Information) (Rimet et al. 2016), and such strain information was also added to the present study when relevant. Altogether, *rbcL* barcodes were available for 66 strains in the TCC collection. The RBGE barcode and strain information has now also been added to the TCC collection. DNA extraction and *rbcL* sequencing of the TCC clones, plus information about the NCBI strains used, are described in Rimet et al. (2016).

Study of morphological characters

RBGE strains: three different analyses of the morphological characters were made for the present study: (1) examination of LM photographs of cleaned material on permanent slides,

(2) a study of the fragile unprepared voucher material dried onto cover slips, especially to see the form of any colonies present, and (3) a scanning electron microscopy (SEM) study of material dried on the 13 mm cover slips, after cleaning the material in situ with 70% HNO₃ and washing with water, as for the LM preparations. Due to time restrictions, detailed studies of voucher material (for colony formation and SEM) were done on a subset of the sequenced isolates (42 strains). For SEM, cover slips were attached to aluminum stubs by carbon discs, coated with platinum, and studied in a LEO Supra 55VP at 5 kV.

TCC strains: LM photographs were not available for all strains and no SEM observations had been made before our study. To obtain material for SEM, all still-living strains were cultured at the Swedish University of Agricultural Sciences, and 23 of these were examined using SEM at RBGE, prepared in the same way as for the RBGE strains. We then studied all available LM photographs of living cultures and cleaned material, and additionally SEM pictures of the 23 strains.

Combined dataset: The selection of strains for SEM was based on a preliminary phylogenetic analysis to analyze diversity, and to cover morphological variability within the clusters, within the constraints of strain availability. Using the morphological characteristics determined with LM and SEM, we then re-examined the strain identifications in relation to published information and, where possible, the morphological characteristics of type material. We also checked commonly used identifications and related literature to study how other authors evaluated this complex. The compilation of morphological features from published species descriptions and type material is available as Table S2.

General remark on morphological data derived from diatom cultures: We are aware that diatom cells in cultures, especially after some time, often decrease in cell size and develop valve deformations, and thus may not necessarily resemble the morphology of natural populations (Mann and Chepurnov 2004). We dealt with this problem by (a) where possible,

using morphological characters derived from LM micrographs taken as soon after isolation as possible where cell length and morphology had not yet changed from the population that existed at the site at the time of sampling, and (b) taking care when interpreting populations with obviously reduced cell sizes coupled to clear deformities during our analyses.

Phylogenetic analyses

The *rbcL* chloroplast gene marker was used to produce a phylogeny of the *Fragilaria* strains. This gene was selected for having 1) available sequences for all our *Fragilaria* strains and b) good efficiency to identify diatoms at species level with DNA metabarcoding when compared to other genes (Kermarrec et al. 2013, Kermarrec et al. 2014). All *rbcL* sequences of the 194 selected strains from the RBGE and TCC collections were aligned using CLUSTAL W Multiple alignment (Thompson et al. 1994) in BioEdit (Hall 1999). The lengths of the Sanger sequences were 1440 base-pairs (bp) for the RBGE strains and at least 574 bp for the TCC strains. To establish a phylogenetic tree, we used a subset of the aligned part trimmed to 1087 bp. The best substitution model was identified with PhyML (Guindon et al. 2010). A first phylogenetic tree was then calculated with RAxML v.8.2.10 (Stamatakis 2014) with the substitution model GTR+I+G. The tree was rooted with *Ulnaria* (30 strains), a neighbor genus of *Fragilaria* (154 strains). To assess branch support, 1000 bootstraps were run. We then removed duplicate sequences, leaving 45 *Fragilaria* and 12 *Ulnaria* sequences. We recalculated the phylogenetic tree, using the same analysis as before. We used a bootstrap threshold of 69% (Soltis and Soltis 2003) to define well-supported groups and thus help define MOTUs. A distance matrix of all long *rbcL* sequences (length > 1086; 178 strains) was calculated by MEGA7 (Kumar et al. 2016) to analyze the number of substitutions per base pair within well-supported phylogenetic clusters.

Material and data accessibility

All material associated with the RBGE strains is stored at the Royal Botanic Garden Edinburgh (for voucher slide accession numbers, see Table S1). All material associated with the TCC strains is accessible through the Thonon Culture Collection (Rimet et al. 2018a). All metadata of both the RBGE and the TCC strains are stored in the open-access R-Syst::diatom reference database; a detailed description of this database and its management is given in (Rimet et al. 2016).

Results

Overall, the phylogenetic tree based on the *rbcL* barcode and rooted with *Ulnaria* (Fig. 1) shows two groups of *Fragilaria* taxa: (1) a modestly supported clade with relatively few sub-branches (the FGRA clade), of which one subgroup was however well-supported; and (2) a very well supported but more heterogeneous clade containing some well supported subgroups together with other strains whose interrelationships are unclear. Below we define and characterize the *Fragilaria* clades that receive good support in the *rbcL* tree. An overview of the morphological characters visible in light and scanning electron microscopy respectively, and a summary of the molecular separations, can be found in Tables 1 and 2. To simplify reading, we used working name abbreviations for the *Fragilaria* groups. Thus, FCAP, FGRA, FPEM, FTEN, FTNS, FVAU, FCRNAPA refer to clones that were assigned, in preliminary identifications, to *F. capucina*, *F. gracilis*, *F. perminuta* (Grunow) Lange-Bert. by Tuji & Williams, *F. tenera* (W. Sm.) Lange-Bert., *F. tenuistriata* Østrup, *F. vaucheriae*, and the group of *F. crotonensis* Kitton-*F. nanoides* Lange-Bert.-*F. pararumpens* Lange-Bert., G. Hofmann & Werum, respectively.

271 ***The FGRA clade***

272 The modestly supported FGRA clade included 77 strains in total, among which there was a
273 well-supported (97%) subgroup, a clade, with 52 strains. Both groups contain many strains
274 from UK, but also from other parts of Europe. 19 of the 77 strains were studied in SEM.
275 Three of the TCC cultures had turned into dwarf forms at the time of study; therefore their
276 morphological characters were considered with care. We first summarize the well-supported
277 clade **FGRA2** separately from the rest of the strains, which form a paraphyletic grade and are
278 referred to as **FGRA1**. All of the strains with an original identification as “*F. gracilis*” fell
279 into the **FGRA** clade.

280 **FGRA2** (52 strains, Fig 2a). 50 of the strains were from UK, 2 from Italy. Two haplotypes
281 were recorded. *Morphology*. No aggregations of cells were observed. Valves mostly linear or
282 linear-lanceolate, gradually narrowing towards bluntly rounded to weakly protracted or
283 subcapitate ends. Length 12–45 μm , breadth 2–3.1 μm at the widest point. The central area is,
284 in most cases, approximately square and the axial area is narrow and linear; sometimes the
285 central area extends into a widened axial area, producing a rhombic shape. Striae always
286 opposite. Stria density 20–24 per 10 μm (mean 22), no areolae visible in light microscope.
287 SEM revealed a rimoportula on the valve face at one of the poles. Two oval to rectangular
288 apical pore fields with 4–8 columns of pores, containing a maximum of four pores each.
289 Spines absent. *Sequence distances*. The sequence distances within the group of FGRA2 of the
290 studied part of *rbcL* of 1087 bp were 0–0.001 substitutions per site.

291 **FGRA1** (25 strains, Fig. 2b). 18 of the strains were from UK, the rest from Italy, Portugal,
292 Luxembourg and Sweden. Five haplotypes were included in this group. *Morphology*. No
293 aggregation of cells was observed. Valves mostly linear, a few strains had linear-lanceolate
294 outlines, gradually narrowing towards bluntly rounded to weakly protracted or subcapitate
295 ends. Length 16–42 μm , breadth 1.8–2.7 μm at the widest point. The central area is, in most

cases, approximately square and the axial area is narrow and linear; sometimes central area extends into a widened axial area, producing a rhombic shape. Striae always opposite, 18–23 per 10 μm (mean 20), no areolae visible in light microscope. SEM revealed a rimoportula on the valve face at one of the poles. Two apical pore fields with oval to rectangular form; 4–8 pore columns with up to four pores each. Spines absent. *Sequence distances*. The sequence distances within the group of FRGA2 of the studied part of *rbcL* of 1087 bp were 0–0.004 substitutions per site.

The heterogeneous Fragilaria clade, containing the FVAU, FCAP1, FCAP2, FTNS, FTEN1, FTEN2 and FPEM subgroups and the residual FCRNAPA strains

All other strains identified as *Fragilaria* species fell into the well-supported (98%) heterogeneous clade. Whereas all strains in the **FGRA** clade had opposite striae, all those in this clade had alternate striae.

Although the deeper relationships in this clade were unclear, it did contain well-supported subgroups. One clade with an identical sequence of the studied part of the *rbcL* barcode (hereafter called **FVAU**) contained several strains originally identified as *F. vaucheriae*. Two other clades contained many strains original identified as *F. capucina* (hereafter called **FCAP1**, 79% support and **FCAP2**, 84% support). Other groups were a clade containing strains all originally identified as *F. perminuta* (**FPEM**, 87% support), and a clade resembling *F. tenuistriata* (hereafter called **FTNS**, 79% support). We also found two clades containing strains with longer valves, resembling *F. tenera* (hereafter called **FTEN1**, 100%, and **FTEN2**, 79% support). There were also many additional strains in the heterogeneous **FCRNAPA** clade, which were irregularly distributed in the phylogenetic tree but still relatively closely

related, according to their sequences. Many of these strains were originally identified as *F. crotonensis*, *F. nanoides* or *F. pararumpens*.

FVAU (6 strains, Fig. 3). The clade was made up of strains from UK (1 strain), the Ile de La Réunion (3 strains), Luxembourg (1 strain) and Italy (1 strain), all strains with identical sequences regarding the studied part of *rbcL*. *Morphology*. LM pictures were available for four strains; these showed long ribbon-band colonies in three, but not in 041SynPO4; instead, cells were found in irregular dense clumps some of them loosely attached to each other. For further details on morphology, see Box 1.

FCAP1 (4 strains, Fig. 4). The clade was made up of one strain from Italy (haplotype 1) and three strains from UK (haplotype 2). *Morphology*. LM and SEM pictures were available for the UK strains. None of the strains formed ribbon-bands, but instead formed loosely irregular aggregates of about ten cells. However, in two strains with elongated cells (~37 µm) most cells came in pairs connected either in the middle or via the entire side. For further details on morphology see Box 2. *Sequence distances*. The sequence distances of the studied part of *rbcL* of 1087 bp were 0–0.003 substitutions per site.

FCAP2 (7 strains, Fig. 5). The clade was made up of two strains from Sweden with identical sequences and five strains from UK, with sequences that were identical to each other but different from the Swedish strains. *Morphology*. LM and SEM pictures were available for all strains. No clear pattern could be observed for colony formation within the clade. One of the Swedish strains formed long ribbon-like bands (>10 cells) relatively loosely connected in the central part of the cell, leaving the ends free. The other Swedish strain was found at most in pairs, some of which were connected by their ends to form irregular stellate colonies. Of the Scottish strains, three were found to form long bands (> 10 cells), one formed shorter bands (2–5 cells), and one occurred mostly as single cells with some pairs. All strains were also found to build irregular aggregates connected by the cell ends. The presence of spines was not

344 correlated with the presence of colonies. For further details on morphology see Box 3.
345 *Sequence distances.* The sequence distances within this group for the studied part of *rbcL* of
346 1087 bp was 0–0.002 substitutions per site.
347

Box 1. *Fragilaria agnesiae* Kahlert & Rimet (Fig.3) (operational name for this article: FVAU)

Description: Valves mostly linear-lanceolate, becoming lanceolate in shorter cells, ends rostrate. Length of the studied strains 9–21 μm (to 65 μm in recently expanded post-auxospore cells). Cell width 4.0–5.4 μm at the center. Central area always unilateral, often strongly so, with a rimmed swelling on the outer valve side. Striae always alternate, 14–16 per 10 μm , with no areolae visible with LM. Most studied strains form long ribbon-like colonies, but shorter bands and irregular loose aggregates also occur. SEM characters: Rimoportula positioned on the valve face at one of the poles. Two apical pore fields with rectangular form, with 8–10 columns of pore columns containing a maximum of five pores each. Spines relatively small, with a round base ending in a round tip. No internal rimmed depression observed in the central area.

Holotype: Material of clone 041SynP04, as preserved on slide BC0041 (Royal Botanic Garden Edinburgh (herbarium abbreviation = E)) illustrated in Fig. 3c, e–i, barcoded by *rbcL* in GenBank accession XXXX, and with preserved DNA at E as EDNA13-0031229.

Authenticated material: Seven isolated strains form the basis of the species description: 041SynP04 (type strain), TCC541, TCC553, TCC558, TCC662, TCC681, TCC547. The TCC strains are all registered and conserved in the Thonon Culture Collection of the INRA at Thonon-les-Bains, France. All strains belong to one haplotype with respect to an identical part of the *rbcL* barcode sequence (1087 bp long).

Type locality: Streams, Pentland Hills, Green Cleuch, above Balerno, Midlothian (UK), sampled 19 May 2012 by David Mann, coordinates: 55°84'42.2"N, 3°31'10.9"W.

Name registration: <http://phycobank.org/100270>

Etymology: The specific epithet refers to the wife of one of the authors.

Similar taxa: This taxon resembles especially *F. rinoi* Almeida et C. Delgado, but this species is described as having no spines and a solitary habit. Other similar species matching in length and form are *F. vaucheriae*, *F. neointermedia* Tuji & Williams, *F. capucina*, *F. pectinalis* and *F. uliginosa* Kulikovskiy, Lange-Bert., Witkowski & Dorofeyuk. However, *F. vaucheriae*, which is the closest in width, should not have more than 14 striae per 10 μm and not form ribbon-like colonies, *F. pectinalis* and *F. uliginosa* also do not form colonies, the latter also has large spatulate spines. Other taxa matching *F. agnesiae* in length and striae density differ from it in the possession of prominent capitate heads (e.g. *F. recapitellata* Lange-Bert. et Metzeltin), or because they are thinner (e.g. *F. perminuta* and *F. microvaucheriae*).

Ecology: The cluster of strains used to describe this new species was isolated from streams in UK (Pentland Hills, Green Cleuch, above Balerno, Midlothian), the Ile de La Réunion (rivière de Bras Caverne, rivière de Langevin – grand Galet, and rivière de Langevin, site amont prise EDF), Luxembourg (river Our at Vianden), and from Italy (Trentino, rivière de Regnana à Amont de Bedollo). There are no direct measurements of water chemistry for any of the collection places, but there is some information for the rivière de Langevin on Ile de La Réunion: pH was 8–8.4, conductivity 8.4–9.3 mS m^{-1} , and oxygen saturation was 113% at a temperature of 21–22 $^{\circ}\text{C}$ (measured 2006 and 2007). The dominant diatom taxa indicated moderately high nutrient concentrations at all three sampling sites, and a possible impact of organic pollution at the rivière de Langevin site amont prise EDF. *F. agnesiae* was recorded from this site with 0.7% in November 2006. The River Our at Vianden, Luxembourg is underlain by schist geology and has good water quality.

Box 2. *Fragilaria heatherae* Kahlert & M. G. Kelly (Fig. 4) (operational name for this article: FCAP1)

Description: Long valves are spindle-shaped with subcapitate ends; short valves are lanceolate with acute to rounded ends. Length of the studied strains 9–38 µm. Cell width 3.3–3.7 at the center. Central area of long cells weakly asymmetric and blurred, whilst, in short cells it is strongly unilateral and clearly separated from the axial area, with a rimmed swelling on the external valve side. Striae always alternate, no areolae visible with LM. Stria density 16 per 10 µm in long cells, 18 per 10 µm in short cells. Cells found in loosely irregular aggregates of about ten cells. SEM characters: Rimoportula positioned on the valve face at one of the poles. Two apical pore fields with rectangular form, with 9 (short cells) or 12–14 (long cells) columns of pores containing a maximum of six pores each. Spines absent.

Holotype: Material of strain 513FraK01 as preserved on slide BC0513 (E), illustrated in Fig. 4a, d, h, barcoded by *rbcL* in GenBank accession XXXX, and with preserved DNA at (E) as EDNA13-0031453.

Authenticated material: Four strains are the basis for the species description: 513FraK01 (type strain), 514FraK01 (in E as slide BC0514), 621FraP11 (in E as slide BC0621), and TCC682 (registered and conserved in the Thonon Culture Collection of INRA at Thonon-les-Bains, France). The RBGE strains belong to one haplotype with respect to an identical part of the *rbcL* barcode sequence (1087 bp long), the TCC strain to a second haplotype.

Type locality: Euden Beck (England, UK), sampled 20 June 2012 by Martyn Kelly, coordinates: 54°66'50.4"N, 1°89'76.5"W.

Name registration: <http://phycobank.org/100271>

Etymology: The specific epithet refers to the wife of one of the authors.

Similar taxa: Size and form resemble *F. capucina* sensu stricto Desm., but this species has spathulate spines in contrast to *F. heatherae* where spines are absent. The size and shape resemble also *Fragilaria pectinalis* (O. F. Müll.) Lyngb. and *Fragilaria microvaucheriae* C. E. Wetzel et Ector; however, the upper limit for the length:width ratio, which provides clear diagnostic characters for these two species, is lower (<8) than for *F. heatherae* (up to 11, wholly including the ranges of the other two). Moreover, the subcapitate form of the ends separates *F. heatherae* from *F. microvaucheriae* (rostrate ends). *F. perminuta* has a higher striae density (17–19), *F. vaucheriae* a lower striae density (9–14).

Ecology: The cluster of strains used to describe this new species was isolated from streams in UK (Euden Beck and River Tay, Pitlochry, Perth & Kinross) and Italy (Trentino rivièr de Regnana à Amont de Bedollo). Euden Beck drains a largely forested catchment with some moorland and rough grazing in the upper catchment. The water is relatively soft and low in nutrients (mean chemistry for June 2011 to June 2012: pH: 7.4; alkalinity: $18 \text{ mg L}^{-1} \text{ CaCO}_3$; conductivity: 10.4 mS m^{-1} ; ammonia-N: 0.02 mg L^{-1} ; nitrate-N: 0.22 mg L^{-1} ; molybdate-reactive P: 0.015 mg L^{-1}). Long-term averages for the River Tay at Aberfeldy also show circumneutral, soft and low nutrient conditions (pH: 7.3, conductivity: 5.3 mS m^{-1} , alkalinity: $11.7 \text{ mg L}^{-1} \text{ CaCO}_3$, $\text{NO}_3\text{-N}$: 0.103 mg L^{-1} , ammonia-N: 0.024 mg L^{-1} , reactive P: 0.013 mg L^{-1} , total P: 0.031 mg L^{-1}).

Box 3. *Fragilaria joachimii* Kahlert (Fig. 5) (operational name for this article: FCAP2)

Description: Long cells: linear; medium-sized cells: lanceolate to spindle-shaped; short cells: lanceolate. Length of the studied strains 5–34 μm . Cell width 3.3–4.6 measured at the center. Central area variable, mostly rectangular, or unilateral with a rimmed swelling. Striae always alternate, 14–16 per 10 μm (up to 19 in some short cells); no areolae visible in LM. No regular pattern of colony formation: cells found in long ribbon bands, or pairs and singular cells, all forming loosely connected aggregates. SEM characters: rimoportula positioned on the valve face at one of the poles. In one cell two rimoportulae were found, one at each pole of one valve. Two apical pore fields with rectangular form, with 5–14 columns of pores containing a maximum of five pores each. A central rimmed depression mostly present in various grades. Very tiny spines irregularly arranged on the valve edges and ends, no spines found on short valves. The presence of spines was not correlated with the presence of long ribbon-like colonies.

Holotype: Material of strain TCC887 (Thonon Culture Collection of the INRA at Thonon–les–Bains, France), illustrated in Fig. 5b, b2, c, j, m, o and p, barcoded by *rbcL* in GenBank accession XXXX.

Name registration: <http://phycobank.org/100272>

Authenticated strains: Seven isolated strains provide the basis of the species description: 042SynP04, 046SynP04, 054SynP04, 171FraB05, 435FraT01 (all in E, as slides BC0042, BC0046, BC0054, BC0171 and BC0435, respectively), and TCC877 and TCC887 (type strain) (registered and conserved in the Thonon Culture Collection of the INRA at Thonon–les–Bains, France). The five RBGE strains belong to one haplotype with respect to an identical part of the *rbcL* barcode sequence (1087 bp long), the two TCC strains to a second haplotype.

Type locality: Broströmmen near Norrtälje city, Sweden, sampled 24 September 2013 by Maria Kahlert, coordinates: 59°75'72.8"N, 18°72'06.0"E.

Etymology: The specific epithet refers to the husband of the author.

Similar taxa: In size and form, *F. joachimii* resembles *F. capucina* Desm. sensu stricto, but *F. capucina* has spatulate spines, in contrast to *F. joachimii*, where the spines are tiny and irregular. The size and form also resemble *F. pectinalis* and *F. microvaucheriae*; however, the length:width ratio (defined as clear diagnostic character for these two species) of *F. joachimii*, is wholly including the ranges of both those species. Moreover, the presence of spines, the formation of colonies and the subcapitate form of the ends separate *F. joachimii* from *F. pectinalis* (no spines or colonies) and *F. microvaucheriae* (rostrate ends). *F. perminuta* has a higher striae density (17–19 per 10 μm), *F. vaucheriae* a lower striae density (9–14 per 10 μm).

Ecology: The cluster of strains used to describe this new species was isolated from streams in Sweden (Norrtäljeån and Broströmmen, both close to Norrtälje city) and Scotland, UK (Pentland Hills, Green Cleuch, above Balerno, Midlothian; Allt a'Bhalachain, Argyll & Bute; and the River Tay, near Aberfeldy, Perth & Kinross). There is a detailed monitoring program for water quality available for the Swedish streams. Both catchments are covered by approx. 30–50% forest, and 20–40% agricultural land. Diatoms have generated IPS values of 13.1–15.8 over the course of several years. Mean pH in Broströmmen was 7.7, mean total phosphorus (TP) 0.046 mg l^{-1} and mean total nitrogen (TN) 1.162 mg l^{-1} . Mean Total Organic Carbon (TOC) was 12.7 mg l^{-1} and mean conductivity 36 mS m^{-1} (measured 2006–2008). Annual mean oxygen saturation was 78% at a mean temperature of 8.0°C (measured 2016). In Norrtäljeån, annual mean temperature was 8.0°C, pH 7.8, conductivity 37 mS m^{-1} , TOC 13 mg m^{-1} , oxygen saturation 83%, TP 0.049 mg l^{-1} and TN 1.666 mg l^{-1} (measured in 2016).

485 Long-term averages for the River Tay at Aberfeldy: pH: 7.3, conductivity: 5.3 mS m⁻¹,
486 alkalinity: 11.7 mg L⁻¹ CaCO₃, NO₃-N: 0.103 mg L⁻¹, ammonia-N: 0.024 mg L⁻¹, reactive P:
487 0.013 mg L⁻¹, total P: 0.031 mg L⁻¹).

488

FPEM (10 strains, Fig. 6). This well-supported clade of 10 strains (87%) comprised three haplotypes, of which the first was represented by two strains from UK and five from Sweden, the second by two, and the third by one strain from France, respectively. All were originally identified as *F. perminuta*. *Morphology*. LM data were available for all strains and SEM for five of them. All strains were rhombic, with a distinct strongly unilateral central area with a rimmed external swelling and internal depression (horse-shoe like structure). Length: 8–23 μm ; width: 3.2–4.6 μm ; stria density: 18 to 21 per 10 μm . No spines were observed in SEM, and the strains did not form colonies. *Sequence distances*. The sequence distances within the group of FPEM of the studied part of *rbcL* of 1087 bp was 0–0.001 substitutions per site.

FTNS (3 strains, Fig. 7). The next well-supported clade (79%) comprised a mixture of one UK and two TCC strains from Sweden, originally identified as *F. mesolepta* Rabenhorst and *F. tenuistriata*. The UK and one of the TCC strains belonged to one haplotype, the other TCC one to a separate one. *Morphology*. For the RBGE strain and one TCC strain both LM and SEM pictures were available; for the other TCC strain LM pictures only. All three strains formed long ribbon-like colonies, with no obvious separation cells. The strains were 34–40 μm long and 4.1–4.5 μm wide, with 16–17 alternating striae per 10 μm . Valves linear, center very slightly almost not constricted, ends bluntly rounded. In SEM, large spatulate spines were observed. The rimoportula was placed at the mantle face junction. *Sequence distances*. The sequence distances within the FTNS clade of the studied part of *rbcL* of 1087 bp were 0–0.001 substitutions per site. For comparison, we also studied the strain 653FraK08, which was also originally identified as *F. mesolepta*, but did not cluster together with the three FTNS strains. This strain had similar morphological features as FTNS, but a different valve outline (Fig. 7b).

FTEN1 (7 strains, Fig. 8). This clade of seven strains (100%) comprised two haplotypes, all from UK, one represented by six strains, the other by one. Of the seven strains, three were

514 originally identified as *F. tenera* or *F. cf. tenera*, two as *F. pararumpens* and two as *F.*
515 *gracilis*. *Morphology*. LM data were available for all strains and SEM for six of them. The
516 strains were quite similar morphologically, ranging from 40–59 µm long, 2.1–3 µm wide,
517 with 18–20 alternating striae per 10 µm. Most cells were solitary or, at most, loosely
518 aggregated. The valves were linear-lanceolate with slightly convex margins and subcapitate
519 apices. The central area was quite broad and slightly inflated in six out of the seven strains
520 and extended into the axial area due to shortened striae. In SEM, the central area was more
521 difficult to observe and marked by ghost striae in all cases. SEM revealed that all strains had
522 very regularly arranged pyramidal spines, often with a sharply bent tip. Spines were present at
523 the poles as well. *Sequence distances*. The sequence distances within the group of FTEN1 of
524 the studied part of *rbcL* of 1087 bp were 0–0.001 substitutions per site.

525 **FTEN2** (3 strains, Fig. 8). This clade was made of three strains of two haplotypes (79%), one
526 from UK and two from Sweden. The Swedish strains were originally identified as *F. cf.*
527 *nanoides*, the Scottish strain as *F. cf. pararumpens*. *Morphology*. LM data were available for
528 all strains and SEM for two of them. Like the FTEN1 clade, the strains were quite similar
529 morphologically, being 50–85 µm long, 2.4–3 µm wide, with 21 alternating striae per 10 µm.
530 Most cells were solitary or at most in pairs; rarely three to four cells were connected at the
531 ends to form a stellate colony. The valves were spindle-shaped with slightly convex margins
532 and subcapitate to capitate apices. The central area was even broader than in FTEN1, and also
533 slightly inflated in all strains, extended far into the axial area due to shortened striae, creating
534 a rhombic central area. Some cells had a quite long section of the central part of the valve
535 having parallel sides, which then abruptly tapered to the ends. In SEM, the central area was
536 also rhombic, and often marked by ghost striae. Spines were present, and often of pyramidal
537 form. One rimoportula was present on each valve, close to the end. However, in contrast to
538 FTEN1, spines were more irregular with various forms, and were often missing in the “neck”

of the cell. Spines could have both round and rectangular quadratic bases, and the tip, often sharply bent, could also form a leaf-like structure in some cells, or was reduced to a wart.

Sequence distances. The sequence distances within the group of FTEN2 of the studied part of *rbcL* of 1087 bp were 0–0.001 substitutions per site.

FCRNAPA: the residual strains of the heterogeneous clade. The original identifications of these strains as *F. crotonensis*, *F. nanoides* or *F. pararumpens* reflects the fact that all of them had very elongate valves.

Discussion

General. Overall, our phylogenetic tree based on *rbcL* and rooted with *Ulnaria* showed two *Fragilaria* clades well-separated both from each other and from the *Ulnaria* strains. One clade contained all strains identified as *F. gracilis*, whereas the other was a heterogeneous group including several well-supported subgroups. Below we try to unravel species taxonomy and names of the *Fragilaria* groups that received good support in the *rbcL* tree.

The FGRA clade. The original identifications of these strains were in most cases *F. gracilis*, except for some of the TCC strains, which had originally been identified as *F. rumpens*. The original description of *F. gracilis* was by Østrup (1910), who described it as having almost linear valves with subcapitate to slightly attenuated ends, giving a length of 63 µm and a width of 3.6 µm, with 20 striae per 10 µm. However, these length and width measurements cannot be confirmed when measuring the original picture of Østrup (1910, Tab V, Fig. 117). Instead, the original drawing has a length of 43 and a width of 2.1 µm, with 25 striae per 10 µm. The microphotographs from the lectotype slide, coll. Østrup 1342, given in Krammer and Lange-Bertalot (1991) and Tuji (2007), show lengths of 28–54 µm, widths of 2–2.7 µm, with

18–24 striae per 10 μm . Tuji (2007) describes the striae as “being parallel throughout”, with SEM pictures showing opposite striae with some irregular parts where striae are alternate. Lange-Bertalot and Ulrich (2014) define the striae as “opposite or alternating”. Note that the term “parallel” refers to the orientation of the striae to each other, while the terms “alternate/opposite” refers to whether the striae on either side of the sternum are opposite each other or alternate. Both Lange-Bertalot and Ulrich (2014) and Tuji (2007) show that *F. gracilis* has no spines, a single transapically orientated rimoportula per valve, and two apical pore fields.

All studied strains falling into the FGRA clade in this study, both those belonging to the well-supported FGRA2 as well as those that did not (FGRA1) fit the descriptions of the type of *F. gracilis* given by Lange-Bertalot and Ulrich (2014) and Tuji (2007). The main difference between the subgroup FGRA2 and the rest of the strains is a higher density of striae in FGRA2 (a mean of 22 vs. 20 per 10 μm), and a tendency to rather more linear valve outline in FGRA1. However, this presents a rather simplistic view of the situation, as the literature on *F. gracilis* and similar taxa is more complicated than this.

The first problem is that the species diagnosis is not clear. Lange-Bertalot and Ulrich (2014) refer to Hofmann et al. (2011) for a species diagnosis, and also refer to Hofmann et al. (2011) when giving size characters for *F. gracilis* in their Table 2. However, Lange-Bertalot and Ulrich (2014) give a length of 20–45 μm , width of 1.9–2.5 μm and 20–23 striae per 10 μm , whereas Hofmann et al. (2011) have 10–60 μm , 2–3 μm and ~20 striae per 10 μm . Furthermore, Table 2 in Lange-Bertalot and Ulrich (2014) gives quite detailed size descriptions, whereas Hofmann et al. (2011) just gives a few (range of length, width and approximate number of striae), thus it is not completely clear what is meant when Lange-Bertalot and Ulrich (2014) note that their size characters of Table 2 are based on “Hofmann et al. (2011) supplemented by own observations”. Other complications are that both Lange-

Bertalot and Ulrich (2014) and Hofmann et al. (2011) state that the ends are slightly protracted and rounded, whereas both have pictures that include weakly subcapitate forms. Both Østrup (1910) and Tuji (2007) note that some valves can have subcapitate ends. Next, whereas neither Tuji (2007) nor Lange-Bertalot and Ulrich (2014) comment on colony formation, Hofmann et al. (2011) state that this taxon is often found in ribbon-like aggregates or attached in stellate colonies.

The second problem is that *F. gracilis* is addressed as part of a ‘difficult to separate complex’, with Hofmann et al. (2011) listing *F. famelica* (Kütz.) Lange-Bert., *F. rumpens*, *F. tenera* and *F. capucina* as “very similar taxa”, of which *F. tenera* would be so similar that it would be “impossible to separate from” *F. gracilis*. Lange-Bertalot and Ulrich (2014) then describe even more similar-looking new species, namely *F. tenuissima*, *F. saxoplanktonica* and *F. aquaplus*.

However, we reason that our FGRA clade fits the characters of *F. gracilis* and cannot be mistaken for any of the other species mentioned above. First, stria density is higher than described for *F. famelica*, *F. rumpens* and *F. capucina* (Table S2). Second, *F. tenera* has been described as possessing spines (Table S2), whereas none were seen on specimens from our clade. Third, in published LM illustrations of the above-mentioned taxa (see references in Table S2), alternate striae are clearly visible, whereas our clade had opposite striae throughout. The separation of FGRA from *F. saxoplanktonica* and *F. aquaplus* is more complicated than from the other taxa, since neither *F. saxoplanktonica* nor *F. aquaplus* possess spines, and both have clearly opposite striae, according to Lange-Bertalot and Ulrich (2014). However, *F. saxoplanktonica* is defined as having denser striae (23–28 according to Lange-Bertalot and Ulrich (2014) than *F. gracilis*, and also the valve form is different (fusiform to needle-shaped) to that of *F. gracilis*.

612 More difficult is the separation of *F. aquaplus*. Whereas its size and striae density are similar
613 to that of *F. gracilis* (Table S2), Lange-Bertalot and Ulrich (2014) state that *F. aquaplus* can
614 be distinguished by having a conspicuously narrow axial area and an arrangement of striae
615 which is “opposing throughout”. Actually, using Hofmann et al. (2011) for identification, *F.*
616 *aquaplus* would be identified as *F. nanana* Lange-Bert. sensu Krammer and Lange-Bertalot
617 (1991). Using instead Cantonati et al. (2017), *F. nanana* is synonymized with *F.*
618 *saxoplanktonica*, even if those authors gave a differential diagnosis of these two taxa in
619 Lange-Bertalot and Ulrich (2014). This confusion is caused by the fact that the name “*F.*
620 *nanana*” was given to a mixture of two species both present in the type slide, as described in
621 Lange-Bertalot and Ulrich (2014). We recommend therefore that the name “*F. nanana*” is not
622 used anymore. The FGRA strains in our study fit the length description of *F. gracilis* and *F.*
623 *aquaplus*, even if the strains with a width of more than 2.5 µm would not fit into Lange-
624 Bertalot and Ulrich (2014)’s description of those two species anymore. There is no clear
625 picture regarding axial area, striae density and form of the ends. Some of the strains in both
626 FGRA2 and FGRA1 would fit the description of *F. gracilis*, others of *F. aquaplus*, others
627 none at all. We conclude from this that separation of *F. gracilis* and *F. aquaplus* is not
628 supported by the *rbcL* barcode. We recommend that the wider definition of *F. gracilis* is used
629 for the whole cluster, as many strains are not identifiable to either species sensu Lange-
630 Bertalot and Ulrich (2014), whereas all of them fit the characters of the type material of
631 *F. gracilis*. Our results support a synonymization of *F. aquaplus* back into the wider concept
632 of *F. gracilis* Østrup sensu Østrup. We consider that *F. gracilis*, if considered in this way, is
633 no longer a part of a ‘difficult to separate complex’, but can be separated in LM from the
634 members of the heterogeneous *Fragilaria* clade using the clear opposite arrangement of the
635 striae, which gives the impression of regularly arranged parallel lines across the valve in LM.
636 The heterogeneous *Fragilaria* clade has alternately arranged striae instead. We do not rule out

the option that the FGRA2 might represent a cryptic species, but if so, it will need a new, different separation than that proposed by Lange-Bertalot and Ulrich for distinguishing between *F. gracilis* and *F. aquaplus*.

The heterogeneous Fragilaria clade, containing the FVAU, FCAP1, FCAP2, FTNS, FTEN1, FTEN2 and FPEM subgroups and the residual FCRNAPA strains

FVAU. The original identification of these strains was *F. vaucheriae* in the UK barcode strain and four TCC strains, and *F. rumpens* in two TCC strains. Wetzel and Ector (2015) used the original type material “Kützing 185” of *F. vaucheriae* to analyze this taxon in detail, including girdle views and SEM pictures. Wetzel and Ector (2015) also noted that the name was first published in Kützing’s exsiccata set Algarum Aquae Dulcis Germanicarum (Decas III, No. 24), albeit without a description or figure, which were added later by Kützing (1833b, p. 560, fig. 38). Less detailed analyses of the original material are also given by Tuji and Williams (2006a) Tuji and Williams (2013), Lange-Bertalot (1980), pl. 4, figs 82–94, 97–102), and Krammer and Lange-Bertalot (1991) (2004, pl. 108, figs 10–15). Wetzel and Ector (2015) also state that the concept for FVAU has been shifting from a broader to a narrower concept. Valves of *F. vaucheriae* type material are linear and narrow with rostrate to subcapitate ends with a unilaterally expanded central area. The length is 14–50 µm, the width 3.8–5.1 µm, and the stria density 11–14 per 10 µm. The striae are described by Wetzel and Ector (2015) in the text as ‘subparallel’, pictures showing an alternate arrangement. For SEM, Tuji and Williams (2013) noted one rimoportula per valve near the poles, large rectangular apical pore fields, and no or very small spines that do not link to sibling cells. Wetzel and Ector (2015) confirm these observations (their figs. 39–53), adding that the sibling cells can remain loosely attached to each other (their figs. 44, 47) but never more than two cells could be found together as observed by Petersen (1938) (e.g. their figs. 6, 18, 44 and 47).

662 Descriptions of *F. vaucheriae* as having linking spines and forming long ribbon-like colonies
 663 (e.g. in Tuji and Williams 2006a) are most probably an error originating from the confusing
 664 and shifting concepts of this species. Wetzel and Ector (2015) and Tuji and Williams (2013)
 665 both refer to Petersen (1938)'s comments on problems with the *F. vaucheriae* type slides. The
 666 taxon identified as *F. intermedia* (Grun.) Grun. in Van Heurck's (1881) Synopsis des
 667 diatomées de Belgique, pl. 45. fig. 11 (but not figs. 9 or 10) and present on slide 552 in
 668 Grunow's collection in Vienna agrees with the original illustration given in Grunow (1860),
 669 and should be considered as synonymous with *F. vaucheriae*, having no or very small spines
 670 and forming no colonies. On the other hand, the individuals from Grunow's slide 31,
 671 identified as *F. intermedia* sensu Grunow in Van Heurck (1881) and illustrated in Van
 672 Heurck's plate 45, figs. 9 and 10 (not fig. 11) have spines and form ribbon-band colonies and
 673 have been described as a new species, *F. neointermedia* Tuji et D.M. Williams (Tuji and
 674 Williams 2013).

675 The characters we found for the five strains with LM pictures available, and the one with
 676 SEM pictures (041SynPO4), do not, unfortunately, yield a single consistent diagnostic
 677 character for the FVAU clade, even within the same haplotype (for a summary of
 678 morphological characters of similar species, see Table S2). The form (linear-lanceolate with \pm
 679 rostrate ends and \pm unilateral central area), length 9-21 μm (to 65 μ in recently expanded post-
 680 auxospore cells), width (4.0–5.4 μm) and striae density (14–16 per 10 μm) of the observed
 681 specimens match only the recently described *F. rinoi* (Delgado et al. 2016). However, *F. rinoi*
 682 was described as having no spines at all, whereas we could frequently observe small spines in
 683 SEM. Furthermore, *F. rinoi* was described as solitary, whereas four of the strains in our study
 684 formed long ribbon-formed colonies. Other species matching FVAU in length and form (*F.*
 685 *vaucheriae*, *F. neointermedia*, *F. capucina*, *F. pectinalis* and *F. uliginosa*) are less than 5.1
 686 μm wide, whereas our specimens were up to 5.4 μm wide. *F. vaucheriae* is closest in width

and form to the specimens we have observed; however, this species should not have more than 14 striae per 10 μm (the lower limit for our specimens). Moreover, whereas the tiny conical spines observed in SEM in 041SynPO4 plus the lack of long ribbon-like colonies in this strain would match both *F. vaucheriae* and *F. pectinalis*, the long ribbon-like colonies of the other strains would exclude those two species. Instead, the colony-formation of these strains, combined with their form, length and stria density, matches *F. neointermedia* and *F. capucina*. As we do not have SEM pictures for the ribbon-forming strains, we are not able to tell something about the rimoportulae (two per valve would fit *F. capucina* only), or the presence of spines (both *F. neointermedia* and *F. capucina* should have linking spines). Other taxa matching in length and striae density fit neither, not *F. recapitellata* (because it has very prominent capitate heads), nor *F. perminuta* and *F. microvaucheriae* (because they should be $< 4 \mu\text{m}$ wide, and most of our valves were at least $4 \mu\text{m}$). In conclusion, no existing species fits the well-supported clade based on the *rbcL* barcode. Moreover, both morphological and molecular analysis of this taxon group will lead to inconsistencies in species identification and links to ecology if no attempt is made to name the well-supported groups. Therefore we describe here the new species *Fragilaria agnesiae* (Box 1) and also suggest that further studies of morphological plasticity of the “real” *F. vaucheriae*, and its phylogenetic relationship to *F. agnesiae* are needed, because one possibility is that *F. vaucheriae* might not form ribbon-like colonies under all circumstances.

FCAP. The original identification of strains in FCAP1 and FCAP2 was either *F. capucina* or *F. rumpens* (TCC) or “*F. capucina* cf. var. *capucina*”, *F. vaucheriae* or “*F. cf. vaucheriae*” (RBGE). One strain was named “*F. cf. gracilis*”. *F. capucina* is a taxon that still is not very well understood, even though its type material has been described at least three times (Krammer and Lange-Bertalot 1991, Tuji and Williams 2006b, Delgado et al. 2015). One difficulty is that the type material includes three different diatoms, which were defined as

712 morphological variations of *F. capucina* by Krammer and Lange-Bertalot (1991). However,
713 these three forms are actually quite distinct and have been suggested to represent three
714 different species (Tuji and Williams 2006b, Delgado et al. 2015, not naming them though),
715 differing not only in outline, but also in the presence of spines and the formation of colonies.
716 (Table S2). Tuji and Williams (2006b) and Delgado et al. (2015) considered that *F. capucina*
717 sensu stricto is the taxon with linear valves, rectangular central area, rostrate ends, two
718 rimoportulae per valve, linking spines, and ribbon-like colonies, and Tuji and Williams
719 (2006b) designated one of these valves as the lectotype of *F. capucina*.

720 Following this definition of *F. capucina*, we consider that the strains of our FCAP1 and
721 FCAP2 clades must be something else, as none of them had linking spines. FCAP1 and
722 FCAP2 were mainly differentiated by the absence of spines and colonies in FCAP1 (cells
723 were connected, at most, in pairs). FCAP2 has irregular, very small spines in some strains,
724 and forms colonies, either as bands or in some other form, even if these were sometimes
725 rather loose. Taxa without linking spines which would fit the size measurements of our FCAP
726 clusters (length 5–38 μm , width 3.3–4.6 μm , stria density- 14.5–16(–19) per 10 μm) are *F.*
727 *pectinalis* or *F. microvaucheriae* for the six strains up to 3.8 μm in width, and *F. rinoi* for the
728 strain that was wider than 4.2 μm . Both *F. pectinalis* and *F. microvaucheriae* would fit
729 FCAP1. According to Wetzel and Ector (2015), the best character to separate those two
730 species is the length to width ratio, being 2–6 in *F. microvaucheriae* and 6–8 in *F. pectinalis*.
731 Our valves, however, had L:W ratios between 2.5 and 10 or 11 so, again, none of the taxa
732 described in literature fit the strains in FCAP1. L:W ratios for cluster FCAP2 range from 5.3
733 to 9.4 (short ones 1.5–2), again spanning the quoted ranges for both *F. pectinalis* and *F.*
734 *microvaucheriae*, so we are once more not able to give one consistent name for these three
735 strains. In any case, the subcapitate ends of FCAP2 do not fit the descriptions of *F.*
736 *microvaucheriae* or *F. rinoi*, both of which have rostrate ends. *F. pectinalis* also has rostrate

ends as well as a similar outline; however, Wetzel and Ector (2015) did not see spines or colony formation, whereas our specimens had tiny irregular spines, and sometimes formed colonies. The two other taxa in the *F. capucina* type material would also fit our FCAP strains' outline, size and striae density, but no information is available on the presence of spines or colony formation. Even more confusing, we found two rimoportulae per valve on one specimen in one of the strains, the rest having only one rimoportula, indicating that the number of rimoportulae might not be a sufficiently stable character on which to separate species. If this is the case then it is not clear how *F. capucina* can be separated from similar taxa, as all other characters overlap with several others (but not *F. recapitellata* which has prominent capitate ends, *F. perminuta* which has a higher striae density (17–19) or *F. vaucheriae* which has a lower striae density (9–14)). In conclusion, we are unable to give an established species name to our FCAP clusters, not even to the well-defined subgroups, or to most of the strains at all, because one or another character of the described species do not fit, and it is not clear which of the morphological characters should have priority when identifying. Therefore, to move forward based on our morphological data and well-supported clades, we propose the name *Fragilaria heatherae* for the FCAP1 clade and *Fragilaria joachimii* for FCAP2 (Boxes 2, 3). It is, nonetheless, still necessary to do further studies to understand the morphological plasticity of *F. pectinalis* and *F. capucina*, if 'typical' forms can be found, isolated and sequenced, and to determine their phylogenetic relationship to *Fragilaria heatherae* and *Fragilaria joachimii*. It is clear from the current phylogenetic tree based on *rbcL* that taxa defined using morphology as *F. capucina* sensu lato fall into several clusters which could reflect both different species, but also morphological and molecular variability.

The **FP** clade was the only group of taxa beside the FGRA clade where the morphological identity seemed to be clear, reflected in the fact that all strains were originally identified as *F.*

762 *perminuta*. All strains were rhombic, with a clear, strongly unilateral central area with a
763 rimmed external swelling and deep internal depression (horse-shoe like structure). Length,
764 stria density, and in most cases also the width fit well to the definition of *F. perminuta* in the
765 original description by Grunow in Van Heurck 1881). Even though three strains are too broad
766 ($> 4 \mu\text{m}$), we still consider our strains to match descriptions of *F. perminuta* well, in that they
767 lacked spines, did not form colonies and had valves with a rhombic outline along with a very
768 clear rimmed swelling/depression, which was more distinct than in strains from other clades
769 where this was observed (in 621FraP11 in FCAP1, in 435FraT01, 042SynP04, and in many
770 valves of TCC887 in FCAP2, and in 041SynP04 in FVAU).

771 **FTNS**. The strains of this clade were originally named *F. mesolepta* or *F. tenuistriata*. Tuji
772 and Williams (2008) analyzed the type material of *F. mesolepta*, *F. tenuistriata* and *F.*
773 *subconstricta*, and noted that all three taxa had certain diagnostic characters (Table S2):
774 whereas *F. mesolepta* has linear valves with a concave fascia and subcapitate ends, both *F.*
775 *tenuistriata* and *F. subconstricta* have linear valves with a linear fascia and rostrate ends. *F.*
776 *tenuistriata* and *F. subconstricta* in turn can be separated by the fact that *F. tenuistriata* has a
777 rimoportula situated on the sternum on the valve face, whereas *F. subconstricta* (and *F.*
778 *mesolepta*) have their rimoportulae on the mantle–valve face junction. Size, spines
779 (spathulate) and colony-formation (long ribbon-like) are similar among all three taxa. The
780 three strains in the FTNS clade all fit the description of *F. subconstricta*, whereas the single
781 similar strain 653FraK08 that did not cluster with the other three is *F. mesolepta*, due to the
782 valve outline (constricted median part, subcapitate poles) and the rimoportula on the mantle–
783 valve face junction. Tuji and Williams (2008) noted that *F. tenuistriata* and *F. subconstricta*
784 cannot be separated in LM other than by the position of the rimoportula. In conclusion, this
785 well-supported FTNS clade is clearly *F. subconstricta* and the single strain not fitting the

cluster is *F. mesolepta*, because all morphological characters fit these species. More research is needed to understand if the position of the rimoportula is sufficient to separate two species.

Our **FTEN1** clade fits the description of *F. tenera* var. *tenera*, as described by Krammer and Lange-Bertalot (1991), Lange-Bertalot and Ulrich (2014) and Almeida et al. (2016) (Table S2). This nicely clustered clade includes taxa originally assigned different names (*F. tenera*, *F. cf. tenera*, *F. gracilis*, *F. pararumpens*), showing, again that identification of *Fragilaria* using LM can be a challenge, and that we need clearer statements of which characters should be used to separate species. One character obviously is the arrangement of the striae, which is opposite in *F. gracilis*, whereas they are alternate throughout in *F. tenera*. Another character might be the ability to form colonies, as *F. pararumpens* has been described to be typically forming long ribbon-like-colonies, whereas *F. tenera* tends to be present mainly as single cells (occasionally pairs), which can be united at the base to form radiating clusters. Another character of the valve outline of our strains was a slightly swollen central area, whereas the illustrations of Hofmann et al. (2011) show a clearly inflated center. Unfortunately, no SEM pictures of *F. pararumpens* are given in the first description (Hofmann et al. 2011), nor did we find any in the literature, thus we do not have information about the presence or formation of spines. We conclude that our FTEN1 clade has all the morphological characters of *F. tenera*. However, more care is needed when describing species in order to be clear about characters that differentiate species from similar taxa. The most useful morphological characters in the case of *F. tenera* are the alternation of the striae (which separate it from *F. gracilis*) and the valve outline with no inflated center (which distinguishes it from *F. pararumpens*).

FTEN2. This clade was quite similar morphologically to FTEN1, identified as *F. cf. nanoides* and *F. cf. pararumpens* respectively, but it clusters separately. The main differences were a higher striae density (21 per 10 μm), more irregularly arranged and formed spines, and that

some cells were elongated around the central part of the valve. The stria density is too high to match *F. tenera*, instead matching *F. tenuissima* (Table S2). That at least a few cells were connected at the ends would also match this to *F. tenuissima*, defined as “may form loose few-celled aggregates” (Lange-Bertalot and Ulrich 2014). However, FTEN2 cannot be identified as *F. tenuissima*, because we observed only one rimoportula per valve, and not two as defined for *F. tenuissima*. No other long *Fragilaria* species match the characters we observed for FTEN2 (Table S2), so again we here have a clade not matching any currently species described from morphological characters. We suggest to refer to this cluster as “*F. tenuissima* with one rimoportula” until more strains and information are available for further studies.

Finally, the phylogenetic relationships of **the residual strains of the heterogeneous clade FCRNAPA** were not clear, as no groups were well supported. Most strains were originally identified as *F. crotonensis*, *F. nanoides* or *F. pararumpens*, with a few (e.g. strains TCC562, TCC589 and TCC705) originally named *F. capucina* despite not resembling the description of *F. capucina* at all (Tuji and Williams 2008). Our attempt to give correct species names to the **FCRNAPA** strains was hampered on one hand because not all morphological characters matched already described species, but mostly because we were not able to make comparisons with some long-celled *Fragilaria* species, because not all characters are described in the original accounts. In particular, the SEM characteristics of *F. pararumpens* and *F. nanoides*, and information on colony formation in *F. nanoides*, are missing. Nonetheless, most strains originally identified as *F. pararumpens* did not match this taxon because they had a higher stria density than *F. pararumpens*. Despite so many long *Fragilaria* taxa having been described, there is no species description to which many of our strains can be fitted. Consequently there is a need to improve our knowledge of the identity and phylogenetic relationships of the **FCRNAPA** clade by collecting and analyzing more strains and sequences,

but also by including the study of colony formation and SEM analysis in the morphological descriptions. More work is certainly needed to understand species diversity and taxonomic relationships in *Fragilaria*, and this entails consideration of all possible characters, morphological, ecological and molecular.

Conclusion

Species names and barcodes.

Our study indicates that a) some species, defined using morphological criteria were consistent with groupings established using *rbcL* sequences, and we recommend that these continue to be recognized. These are: *F. gracilis* Østrup sensu Østrup (synonym pro parte: *F. aquaplus* Lange-Bertalot and Ulrich), *F. perminuta* (Grunow) Lange-Bertalot by Tuji & Williams, *F. tenera* (W. Smith) Lange-Bertalot, *F. subconstricta* Østrup, b) even for those species, however, the original identification using LM sometimes differed from the final one confirmed by molecular and morphological characters (including those visible only with SEM), confirming that the identification of *Fragilaria* by LM is difficult. However using a single-gene ‘barcode’ approach provides an effective tool to overcome problems. We also conclude that c) more strains are needed to analyze relationships, especially between strains of long *Fragilaria* taxa, including some apparently clearly-defined species such as *F. crotonensis*. Moreover, we highlighted that d) some of our well-supported clades (FCAP1 and FCAP1, FVAU, FTEN2), as well as some of the less well-supported groups include strains that could not be identified using the current morphological literature, either because the morphological characters of the strains in the group did not fit any of the described taxa, or because several taxa would fit and there was not sufficient clarity on differentiating criteria. In order to ensure the recognition and reporting of the well-supported clusters in our reference

database, especially during subsequent future DNA metabarcoding analysis, we decided to describe three new species: *Fragilaria agnesiae* (FVAU), *Fragilaria heatherae* (FCAP1), and *Fragilaria joachimii* (FCAP2). As a help to identify *Fragilaria* in LM, we also provide a summary of published information on *Fragilaria* morphology (Table S2) and a summary of the LM characters of the taxa we have studied, including suggestions how to solve identification problems (Table S3). Finally, we note that e) there are well-defined subgroups within one morphological species (here FGRA), whose geographical distribution and ecology need further study. If we can find meaningful ecological differences between FGRA1 and FGRA2, it would be worthwhile to define these as distinct (albeit cryptic) species that can be separated by DNA sequences more easily than using LM, where only the stria density differs between these groups, and even this character is overlapping.

Ecological assessment. A complete overhaul of the taxonomic mess in *Fragilaria sensu stricto* will take a long time. However, we need to find out which taxa can be pooled or need to be separated in curated databases in order to achieve harmonized taxa lists. Indeed, DNA approaches reinforce the need for curated and operational taxa lists, linked to their DNA barcodes, for both research and environmental monitoring of freshwater ecosystems. Our results show that our well-supported clusters originate in different parts of Europe, supporting the idea that those clades are not regionally restricted in distribution. Especially for biomonitoring purposes, we need to know how differences in a barcode relate to differences in ecology. Thus, the next step is to relate the different clades to differences in environmental variables in order to determine whether the better taxonomic discrimination obtained using DNA metabarcoding translates into better ecological understanding.

In this paper we have used molecular biology, in effect, to test and validate a taxonomy based on sometimes inadequate descriptions of morphological characters. Moving forward, we will continue trying to provide a classification that is operable simultaneously by traditional and

molecular approaches and that allows maximum continuity with what has done before, and improved insights into both taxonomy and ecology of diatom species.

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References

- Almeida, P. D., Morales, E. A., Wetzel, C. E., Ector, L. & Bicudo, D. C. 2016. Two new diatoms in the genus *Fragilaria* Lyngbye (Fragilariophyceae) from tropical reservoirs in Brazil and comparison with type material of *F. tenera*. *Phytotaxa* **246**:163–83.
- Amato, A., Kooistra, W., Ghiron, J. H. L., Mann, D. G., Proschold, T. & Montresor, M. 2007. Reproductive isolation among sympatric cryptic species in marine diatoms. *Protist* **158**:193-207.
- Battarbee, R. W., Juggins, S., Gasse, F., Anderson, N. J., Bennion, H., Cameron, N. G., Ryves, D. B., Pailles, C., Chalief, F. & Telford, R. 2001. *European Diatom Database (EDDI). An Information System for Palaeoenvironmental Reconstruction*. ECRC Research Report No. 81. pp. 94.

- 909 Cantonati, M., Kelly, M. G. & Lange-Bertalot, H. 2017. *Freshwater Benthic Diatoms of*
910 *Central Europe: Over 800 Common Species Used in Ecological Assessment. English*
911 *edition with updated and added species*. Koeltz Botanical Books, Oberreifenberg. pp.
912 942.
- 913 Cemagref 1982. Etude des méthodes biologiques d'appréciation quantitative de la qualité des
914 eaux. Q.E. Lyon-A.F.Bassion Rhône-Méditerranée-Corse, pp. 218.
- 915 Delgado, C., Novais, M. H., Blanco, S. & Almeida, S. F. P. 2015. Examination and
916 comparison of *Fragilaria candidagilae* sp. nov. with type material of *Fragilaria*
917 *recapitellata*, *F. capucina*, *F. perminuta*, *F. intermedia* and *F. neointermedia*
918 (Bacillariophyta, Fragilariaceae). *Phytotaxa* **231**:1-18.
- 919 Delgado, C., Novais, M. H., Blanco, S. & Almeida, S. F. P. 2016. *Fragilaria rinoi* sp nov
920 (Fragilariales, Fragilariophyceae) from periphytic river samples in Central Portugal.
921 *Eur. J. Taxon.* **248**:1-16.
- 922 Evans, K. M., Wortley, A. H., Simpson, G. E., Chepurnov, V. A. & Mann, D. G. 2008. A
923 molecular systematic approach to explore diversity within the *Sellaphora pupula*
924 species complex (Bacillariophyta). *J. Phycol.* **44**:215-31.
- 925 Grunow, A. 1860. Über neue oder ungenügend gekannte Algen. Erste Folge, Diatomeen,
926 Familie Naviculaceen. *Verh. Zool. Bot. Ges. Wien* **10**:503–82.
- 927 Guindon, S., Dufayard, J.-F., Lefort, V., Anisimova, M., Hordijk, W. & Gascuel, O. 2010.
928 New Algorithms and Methods to Estimate Maximum-Likelihood Phylogenies:
929 Assessing the Performance of PhyML 3.0. *Syst. Biol.* **59**:307-21.
- 930 Hall, T. A. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis
931 program for Windows 95/98/NT. *Nucleic Acids Symp. Ser.* **41**:95-98.

- 932 Hofmann, G., Werum, M. & Lange - Bertalot, H. 2011. *Diatomeen im Süßwasser-Benthos*
 933 *von Mitteleuropa. Bestimmungsflora Kieselalgen für die ökologische Praxis. Über 700*
 934 *der häufigsten Arten und ihre Ökologie*. A.R.G.Gantner Verlag K.G., Rugell, pp. 908.
- 935 Kahlert, M. 2011. Framtagande av gemensamt delprogram ”Kiselalger i vattendrag”:
 936 Underlag för utformning av övervakningsprogram och verifiering av kiselalgsindex.
 937 *Länsstyrelsens rapporter*. Karlskrona, pp. 62.
- 938 Kahlert, M., Albert, R.-L., Anttila, E.-L., Bengtsson, R., Bigler, C., Eskola, T., Galman, V.,
 939 Gottschalk, S., Herlitz, E., Jarlman, A., Kasperoviciene, J., Kokocinski, M., Luup, H.,
 940 Miettinen, J., Paunksnyte, I., Piirsoo, K., Quintana, I., Raunio, J., Sandell, B., Simola,
 941 H., Sundberg, I., Vilbaste, S. & Weckstrom, J. 2009. Harmonization is more important
 942 than experience-results of the first Nordic-Baltic diatom intercalibration exercise 2007
 943 (stream monitoring). *J. Appl. Phycol.* **21**:471-82.
- 944 Kahlert, M., Jarlman, A., Sundberg, I. & Herlitz, E. 2017. Taxalista - kiselalger i svenska
 945 sötvatten. Available at:
 946 [http://miljodata.slu.se/mvm/Content/Static/Current/Kiselalger%20i%20svenska%20sö](http://miljodata.slu.se/mvm/Content/Static/Current/Kiselalger%20i%20svenska%20sötvatten.xlsx)
 947 [tvatten.xlsx](http://miljodata.slu.se/mvm/Content/Static/Current/Kiselalger%20i%20svenska%20sötvatten.xlsx) (last accessed 18 July 2018).
- 948 Kelly, M., Boonham, N., Juggins, S., Kille, P., Mann, D., Pass, D., Sapp, M., Sato, S. &
 949 Glover, R. 2018. *A DNA based diatom metabarcoding approach for Water*
 950 *Framework Directive classification of rivers*. SC140024/R, Environment Agency,
 951 Bristol. ISBN: 978-1-84911-406-6, pp. 157.
- 952 Kelly, M. G. & Whitton, B. A. 1995. Trophic diatom index - a new index for monitoring
 953 eutrophication in rivers. *J. Appl. Phycol.* **7**:433-44.
- 954 Kermarrec, L., Franc, A., Rimet, F., Chaumeil, P., Frigerio, J.-M., Humbert, J.-F. & Bouchez,
 955 A. 2014. A next-generation sequencing approach to river biomonitoring using benthic
 956 diatoms. *Freshw. Sci.* **33**:349-63.

- 957 Kermarrec, L., Franc, A., Rimet, F., Chaumeil, P., Humbert, J. F. & Bouchez, A. 2013. Next-
958 generation sequencing to inventory taxonomic diversity in eukaryotic communities: a
959 test for freshwater diatoms. *Mol. Ecol. Res.* **13**:607-19.
- 960 Krammer, K. & Lange-Bertalot, H. 1991. *Bacillariophyceae. 3. Teil: Centrales,*
961 *Fragilariaceae, Eunotiaceae. In* Ettl, H., Gerloff, J., Heynig, H. & Mollenhauer, D.
962 [Eds] *Süßwasserflora von Mitteleuropa*, vol. 2/3. Gustav Fischer Verlag, Stuttgart &
963 New York, pp. 576.
- 964 Kumar, S., Stecher, G. & Tamura, K. 2016. MEGA7: Molecular Evolutionary Genetics
965 Analysis Version 7.0 for Bigger Datasets. *Mol. Biol. Evol.* **33**:1870-74.
- 966 Lange-Bertalot, H. 1980. Zur systematischen Bewertung der bandförmigen Kolonien bei
967 *Navicula* und *Fragilaria*. *Nova Hedwigia* **33**:723-787.
- 968 Lange-Bertalot, H. & Ulrich, S. 2014. Contributions to the taxonomy of needle-shaped
969 *Fragilaria* and *Ulnaria* species. *Lauterbornia* **78**:1-73.
- 970 Larras, F., Montuelle, B. & Bouchez, A. 2013. Assessment of toxicity thresholds in aquatic
971 environments: Does benthic growth of diatoms affect their exposure and sensitivity to
972 herbicides? *Science of The Total Environment* **463-464**:469-77.
- 973 Lecointe, C., Coste, M. & Prygiel, J. 1993. "Omnidia": software for taxonomy, calculation of
974 diatom indices and inventories management. *Hydrobiologia* **269**:509-13.
- 975 Lecointe, M. 2018. Omnidia software version 6.0.6. Omnidia.
- 976 Mann, D. G. & Chepurnov, V. A. 2004. What have the Romans ever done for us? The past
977 and future contribution of culture studies to diatom systematics. *Nova Hedwigia*
978 **79**:237-91.
- 979 Mann, D. G., Sato, S., Trobajo, R., Vanormelingen, P. & Souffreau, C. 2010. DNA barcoding
980 for species identification and discovery in diatoms. *Cryptogamie Algol.* **31**:557-77.

- 981 Morales, E. 2010. *Fragilaria vaucheriae*. In: Diatoms of North America. Available at:
982 https://diatoms.org/species/fragilaria_vaucheriae (last accessed July 17 2018).
- 983 Mougin, C., Artige, E., Marchand, F., Mondy, S., Ratie, C., Sellier, N., Castagnone-Sereno,
984 P., D'Acier, A. C., Esmenjaud, D., Faivre-Primot, C., Granjon, L., Hamelet, V., Lange,
985 F., Pages, S., Rimet, F., Ris, N. & Salle, G. 2018. BRC4Env, a network of Biological
986 Resource Centres for research in environmental and agricultural sciences. *Environ.*
987 *Sci. Poll. Res. Int.* <https://doi.org/10.1007/s11356-018-1973-7>.
- 988 Petersen, J. B. 1938. *Fragilaria intermedia*-*Synedra vaucheriae*. *Bot. Not.* **1938**:164.
- 989 Poulíčková, A., Veselà, J., Neustupa, J. & Skaloud, P. 2010. Pseudocryptic diversity versus
990 cosmopolitanism in diatoms: a case study on *Navicula cryptocephala* Kütz.
991 (Bacillariophyceae) and morphologically similar taxa. *Protist* **161**:353-69.
- 992 Rimet, F. & Bouchez, A. 2012a. Biomonitoring river diatoms: Implications of taxonomic
993 resolution. *Ecol. Ind.* **15**:92-99.
- 994 Rimet, F. & Bouchez, A. 2012b. Life-forms, cell-sizes and ecological guilds of diatoms in
995 European rivers. *Knowl. Managt. Aquatic Ecosyst.* **406**:01.
- 996 Rimet, F., Chardon, C., Lainé, L., Bouchez, A., Jacquet, S., Domaizon, I. & Guillard, J.
997 2018a. Thonon Culture Collection -TCC- a freshwater microalgae collection. Portail
998 Data Inra.
- 999 Rimet, F., Chaumeil, P., Keck, F., Kermarrec, L., Vasselon, V., Kahlert, M., Franc, A. &
1000 Bouchez, A. 2016. R-Syst::diatom: an open-access and curated barcode database for
1001 diatoms and freshwater monitoring. *Database-Oxford* **2016**:1–21.
- 1002 Rimet, F., Vasselon, V., A-Keszte, B. & Bouchez, A. 2018b. Do we similarly assess diversity
1003 with microscopy and high-throughput sequencing? Case of microalgae in lakes.
1004 *Organisms Diversity & Evolution* **18**:51-62.

- 1005 Sarno, D., Kooistra, W., Balzano, S., Hargraves, P. E. & Zingone, A. 2007. Diversity in the
1006 genus *Skeletonema* (Bacillariophyceae): III. Phylogenetic position and morphological
1007 variability of *Skeletonema costatum* and *Skeletonema grevillei*, with the description of
1008 *Skeletonema ardens* sp nov. *J. Phycol.* **43**:156-70.
- 1009 Snoeijs, P. & Potapova, M. e. 1995. *Intercalibration and distribution of diatom species in the*
1010 *Baltic Sea. Volume 3.* Opulus Press, Uppsala, pp. 125.
- 1011 Soltis, P. S. & Soltis, D. E. 2003. Applying the bootstrap in phylogeny reconstruction. *Stat.*
1012 *Sci.* **18**:256-67.
- 1013 Stamatakis, A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of
1014 large phylogenies. *Bioinformatics* **30**:1312-13.
- 1015 Thompson, J. D., Higgins, D. G. & Gibson, T. J. 1994. Clustal-W - improving the sensitivity
1016 of progressive multiple sequence alignment through sequence weighting, position-
1017 specific gap penalties and weight matrix choice. *Nucleic Acids Res.* **22**:4673-80.
- 1018 Tuji, A. 2007. Type examination of *Fragilaria gracilis* Østrup (Bacillariophyceae). *Bull. Nat.*
1019 *Mus. Nat. Sci., ser. B, Botany* **33**:9-12.
- 1020 Tuji, A. & Williams, D. M. 2006a. Examination of the type material of *Synedra rumpens* =
1021 *Fragilaria rumpens*, Bacillariophyceae. *Phycol. Res.* **54**:99-103.
- 1022 Tuji, A. & Williams, D. M. 2006b. Typification of *Conferva pectinalis* O.F. Müll.
1023 (Bacillariophyceae) and the identity of the type of an alleged synonym. *Fragilaria*
1024 *capucina* Desm. *Taxon* **55**:193.
- 1025 Tuji, A. & Williams, D. M. 2008. Examination of type material of *Fragilaria mesolepta*
1026 Rabenhorst and two similar, but distinct, taxa. *Diatom Research* **23**:503-10.
- 1027 Tuji, A. & Williams, D. M. 2013. Examination of types in the *Fragilaria vaucheriae*-
1028 *intermedia* species complex. *Bull. Nat. Mus. Nat. Sci., ser. B, Botany* **39**:1-9.
- 1029 Van Heurck, H. 1881. *Synopsis des diatomées de Belgique. Atlas.* Anvers, pp. 235.

- 1030 Van Dam, H., Mertens, A. & Sinkeldam, J. 1994. A coded checklist and ecological
1031 indicator values of freshwater diatoms from The Netherlands. *Neth. J. Aquat. Ecol.*
1032 **28**:117-33.
- 1033 Vanormelingen, P., Evans, K. M., Chepurnov, V. A., Vyverman, W. & Mann, D. G. 2013.
1034 Molecular species discovery in the diatom *Sellaphora* and its congruence with mating
1035 trials. *Fottea* **13**:133-48.
- 1036 Vasselon, V., Rimet, F., Tapolczai, K. & Bouchez, A. 2017. Assessing ecological status with
1037 diatoms DNA metabarcoding: Scaling-up on a WFD monitoring network (Mayotte
1038 island, France). *Ecol. Ind.* **82**:1-12.
- 1039 Wetzel, C. E. & Ector, L. 2015. Taxonomy and Ecology of *Fragilaria microvaucheriae* sp.
1040 nov. and Comparison with the Type Materials of *F. uliginosa* and *F. vaucheriae*.
1041 *Cryptogamie Algal.* **36**:271-89.
- 1042 Østrup, E. V. 1910. *Danske diatoméer*. C. A. Reitzel, Kjøbenhavn.
- 1043
- 1044
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1046 **Tables**

1047 Table 1. Summary of morphological data for the well-supported clusters of the studied
1048 *Fragilaria* strains in this study. Characters visible in light microscopy.

1049

1050 Table 2. Summary of morphological data for the well-supported clusters of the studied
1051 *Fragilaria* strains in this study. Characters visible in scanning electron microscopy.
1052 Information on molecular phylogeny based on a *rbcL* barcode included.

1053

1054

Figures

Figure legends

Fig. 1 Phylogeny of the studied strains of *Fragilaria*, rooted with strains of the genus *Ulnaria*, on a 1087 bp long part of the *rbcL* barcode. Bootstrap values above 69% are given for each node. Scale bar: number of substitutions per site.

Fig. 2a. *Fragilaria gracilis* Østrup, studied strains of the well-supported cluster FGRA2. (a–e) Light microscopy (LM); example of valve views (a) 166FraB05, (b) 139FraB04, (c) 177FraB05, (d) 148FraB04, (e) 170FraB05; (f–j) Scanning electron microscopy (SEM); (f) general external view of valve of 166FraB05, no spines present, (g) external and (h) internal view of central area of 166FraB05, (i) external and (j) internal view of head poles with pore fields and rimoportula (i) 726FraB12, (j) 166FraB05. Scale bar 10 µm (a–f), or 1 µm (g–j).

Fig. 2b. *Fragilaria gracilis* Østrup, studied strains of the cluster FGRA1. (a–e) Light microscopy (LM); (a) living culture of TCC869, not forming colonies, (b–e) examples of valve views (b) 907FraR05, (c) 185FraB06, (d) TCC869, (e) 005FraP02; (f–k) Scanning electron microscopy (SEM); (f) general external view of valve of 488FraR03, no spines present, (g) general internal view of valve of TCC869, (h) external and (i) internal view of central area of 185FraB06 (h) and 018FraP02, (j) external and (k) internal view of head poles with pore fields and rimoportula of 185FraB06. Scale bar 10 µm (a–g), or 1 µm (h, k).

Fig. 3. *Fragilaria agnesiae* (operational name for this article: FVAU): (a–d) Light microscopy (LM); (a, b) living cultures (a) of TCC541, forming ribbon-band colonies, (b) of TCC541,

1078 valve and girdle view, (c) valve view of 041SynP04, (d) valve of TCC553. (e–i) Scanning
 1079 electron microscopy (SEM) of 041SynP04; (e, f) general external view of valve and girdle
 1080 view, small spines visible, (g) internal view of central area, no rimmed depression visible,
 1081 external (h) and (i) internal view of head pole with transapical rimoportula on valve face and
 1082 pore field visible. Scale bar 10 μm (a–d), 5 μm (e, f), or 1 μm (g–i).

1083

1084 Fig. 4. *Fragilaria heatherae*, studied strains (operational name for this article: FCAP1): (a–c)
 1085 Light microscopy (LM); (a) valve view of 513FraK01, (b) 514FraK01 and (c) 621FraP11. (d–
 1086 h) Scanning electron microscopy (SEM); general external (d, 513FraK01) and internal (e,
 1087 514FraK01) view (no rimmed depression visible in central area), (f) external view of central
 1088 area of 514FraK01, (g) internal view of central area of 621FraP11 with rimmed depression,
 1089 external (h, 513FraK01) and internal (i, 514FraK01) view of head pole with transapical
 1090 rimoportula on valve face and pore field visible. Scale bar 10 μm (a–c), 5 μm (d, e), 1 μm (f–
 1091 i).

1092

1093 Fig. 5. *Fragilaria joachimii*, studied strains (operational name for this article: FCAP2): (a–j)
 1094 Light microscopy (LM); (a), living culture of TCC877, forming ribbon-band colonies, (b)
 1095 living culture of TCC887, forming stellate irregular colonies, (c) valve view of TCC887, (d–j)
 1096 valve view of all strains forming the new described species: (d) 042SynP04, (e) 046SynP04,
 1097 (f) 054SynP04, (g) 171FraB05, (h) 435FraT01 (i) TCC877, (j) TCC887; Scanning electron
 1098 microscopy (SEM); (k) ribbon-band colonies of 042SynP04, (l) general external view of valve
 1099 of 054SynP04, (m) external view of central area of 171FraB05, showing tiny spines, (m2–m5)
 1100 internal views of central area showing various grades of a rimmed depression, (m2)
 1101 435FraT01, (m3) 171FraB05, (m4) TCC877, (m5) 046SynP04, (n–p) detailed view of head

poles with pore fields; (n) external, tiny spines of 046SynP04, (o) external, tiny spines,
rimoportula of TCC887, (p) internal, rimoportula of 054SynP04. Scale bar 10 μm (a–i), 5 μm
(l, m2–m5), or 1 μm (m–p).

1105

Fig. 6. *Fragilaria perminuta* (Grunow) Lange-Bertalot by Tuji & Williams (operational name
for this article: FPEN). (a, b) Light microscopy (LM); (a) living culture of TCC829, not
forming colonies, (b) valve views of 038FraP04; (c–f) Scanning electron microscopy (SEM);
(c) general external valve view of 043SynP04, (d) detailed internal view on central area with
rimmed depression of 038FraP04, (e) external and (f) view of head pole with pore fields and
rimoportula, (e) of TCC 866, (f) of 038FraP04. Scale bar 10 μm (a, b), 5 μm (c), or 1 μm (d–
f).

1113

Fig. 7. *Fragilaria subconstricta* Østrup (operational name for this article: FTNS). (a–e) Light
microscopy (LM): (a) living culture of TCC867, forming long ribbon-band colonies, (b–d)
valve views of (b) 015FraP02, (c) TCC867, (d) TCC868; (e–h) Scanning electron microscopy
(SEM); (f) general external girdle view 015FraP02, (f) detailed external girdle view of head
poles with pore fields and rimoportula on mantle/valve junction (arrow) of 015FraP02, (g)
detailed external valve view of head poles with pore field of 015FraP02, (h) detailed external
valve view of head poles with rimoportula on mantle/valve junction of 015FraP02. Scale bar
20 μm (a), 10 μm (b–d), 5 μm (e), or 1 μm (f–h).

1122

Fig. 7b. *Fragilaria mesolepta* Rabenhorst, studied strain 653FraK08, to compare with
Fragilaria subconstricta Østrup (Fig. 7). (a) Light microscopy (LM), general external view of
valve; (b–d) Scanning electron microscopy (SEM); (b) general external girdle and internal

valve view, (c) external and (d) internal view of head poles with close-up of the spatular spines, pore fields and rimoportula on mantle/valve junction. Scale bar 10 μm (a, b), or 1 μm (c, d).

Fig. 8. *Fragilaria tenera* (W. Smith) Lange-Bertalot (operational name for this article: FTEN1). (a–g) Light microscopy (LM); examples of valve views of (a) 343FraT01, (b) 575FraK01, (c) 613FraP11, (d) 625FraP11, (e) 662FraK09, (f) 726FraB12, (g) 732FraB12; (h–j) Scanning electron microscopy (SEM); (h) general external valve view of 343FraT01, (i) external view of central area of 613FraP11, showing pyramidal spines, (j) external view of head poles with pore fields and rimoportula of 613FraP11. Scale bar 10 μm (a–h), 5 μm (i, j).

Fig. 9. *Fragilaria* sp. 1 (Operational name for this article: FTEN2). (a–f) Light microscopy (LM); (a) living culture of TCC870, not forming colonies, (b–f) examples of valve views of (b–c) TCC870, (d–e) TCC871, (f) 032FraP02; (g–i) Scanning electron microscopy (SEM); (g) general external valve view of TCC871, (h) external view of central area of 032FraP02, showing pyramidal spines, (i) external view of head poles with pore fields and rimoportula of 032FraP02. Scale bar 20 μm (b–e), 10 μm (f, g), or 1 μm (h, i).

1144 *Figures (separate files)*

1145

1146 **Supplementary Tables** (*separate files*)

1147 Table S1. Strain information, including clone identifier, voucher slide identifier, initial and
1148 final taxon identifications and operational name for this article, collection information, *rbcL*
1149 sequences, and GenBank accession numbers (where appropriate).

1150

1151 Table S2. Compiled information on the genus *Fragilaria*, retrieved as correct as possible from
1152 published material. Length, width (at widest point of valve), number of striae per 10µm, and
1153 information on spines (visible in SEM only) and colony formation (visible on uncleaned
1154 material only) is given. More details and photographs can be found directly in the
1155 publications. This table does not claim to be exhaustive. Information taken from ^t text or ^f
1156 figures in the specified references. SEM: scanning electronic microscopy. LM: light
1157 microscopy. no info = no information is given in the reference. RP: rimoportula. Striae p =
1158 opposite and parallel, a = alternating, pa: both alternatively.

1159

1160 Table S3. Towards more harmonized taxa names: Suggestions on how to separate *Fragilaria*
1161 species by morphological characters in the LM (based on type descriptions).

1162

1163 **Supplementary Figure** (*separate file*)

1164

1165 Fig. S1. Phylogeny of the studied strains of *Fragilaria*, rooted with strains of the genus
1166 *Ulnaria*, on a 1087 bp long part of the *rbcL* barcode. Bootstrap values above 69% are given
1167 for each node. Scale bar: number of substitutions per site. Full tree with geographical
1168 information included.

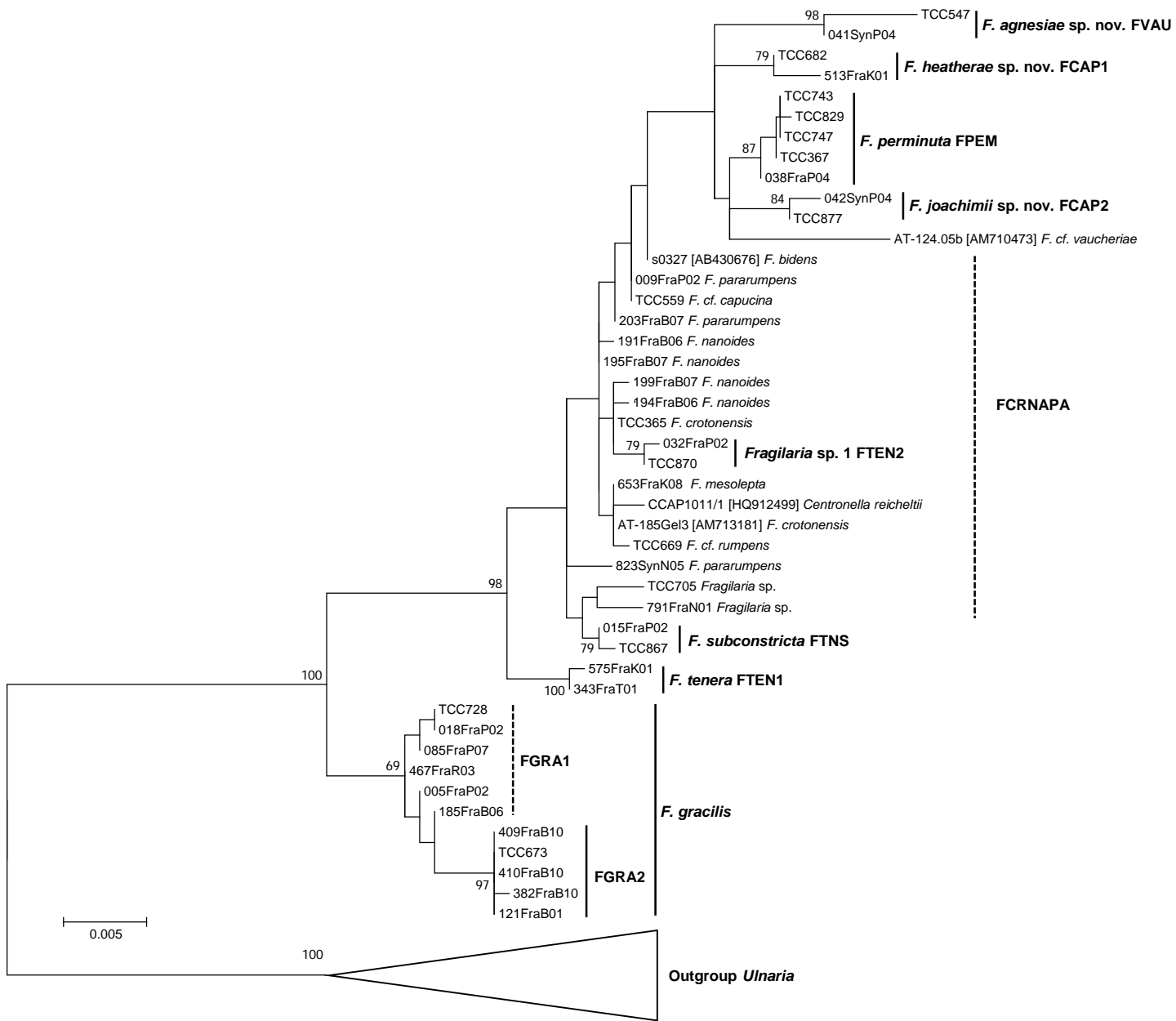
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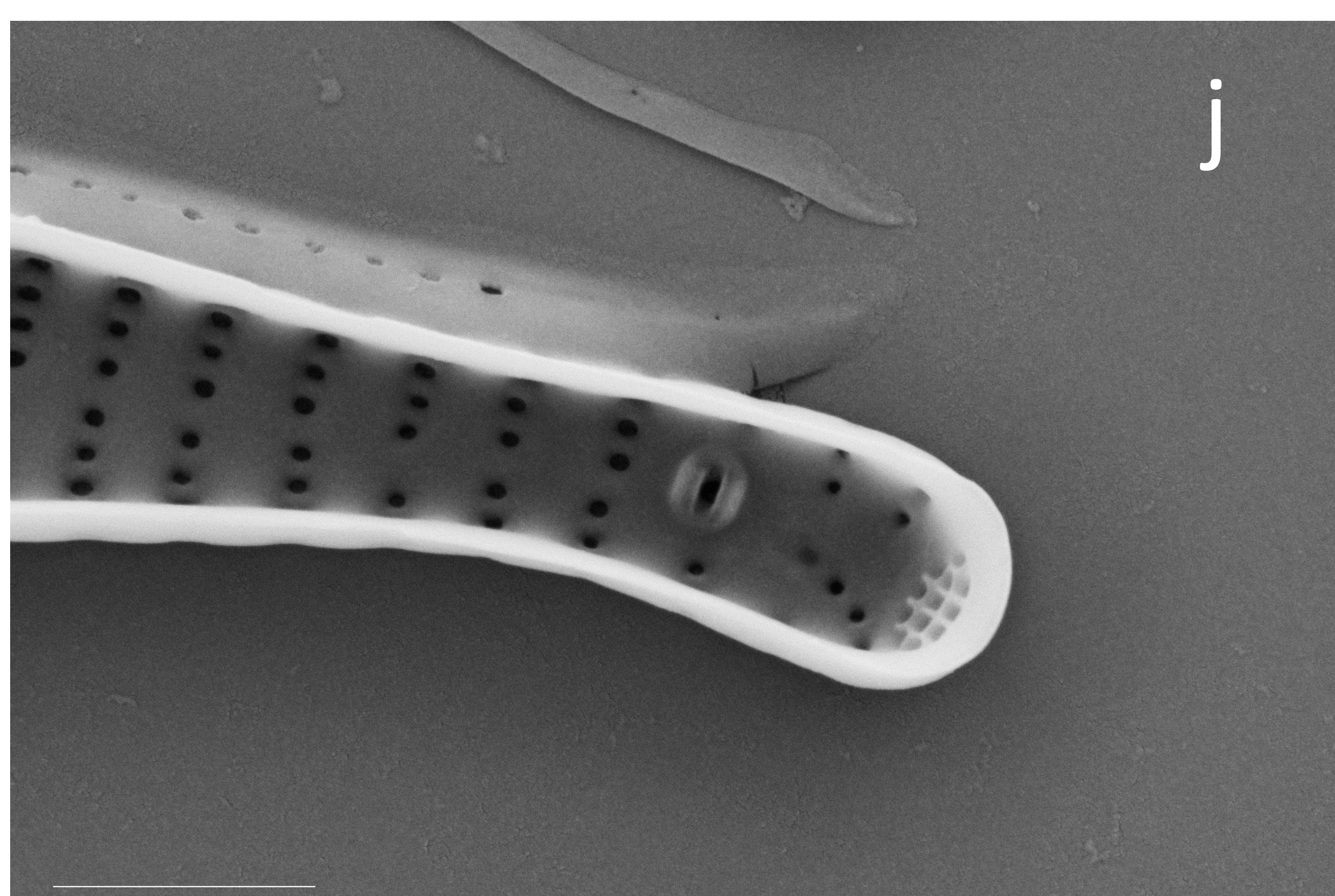
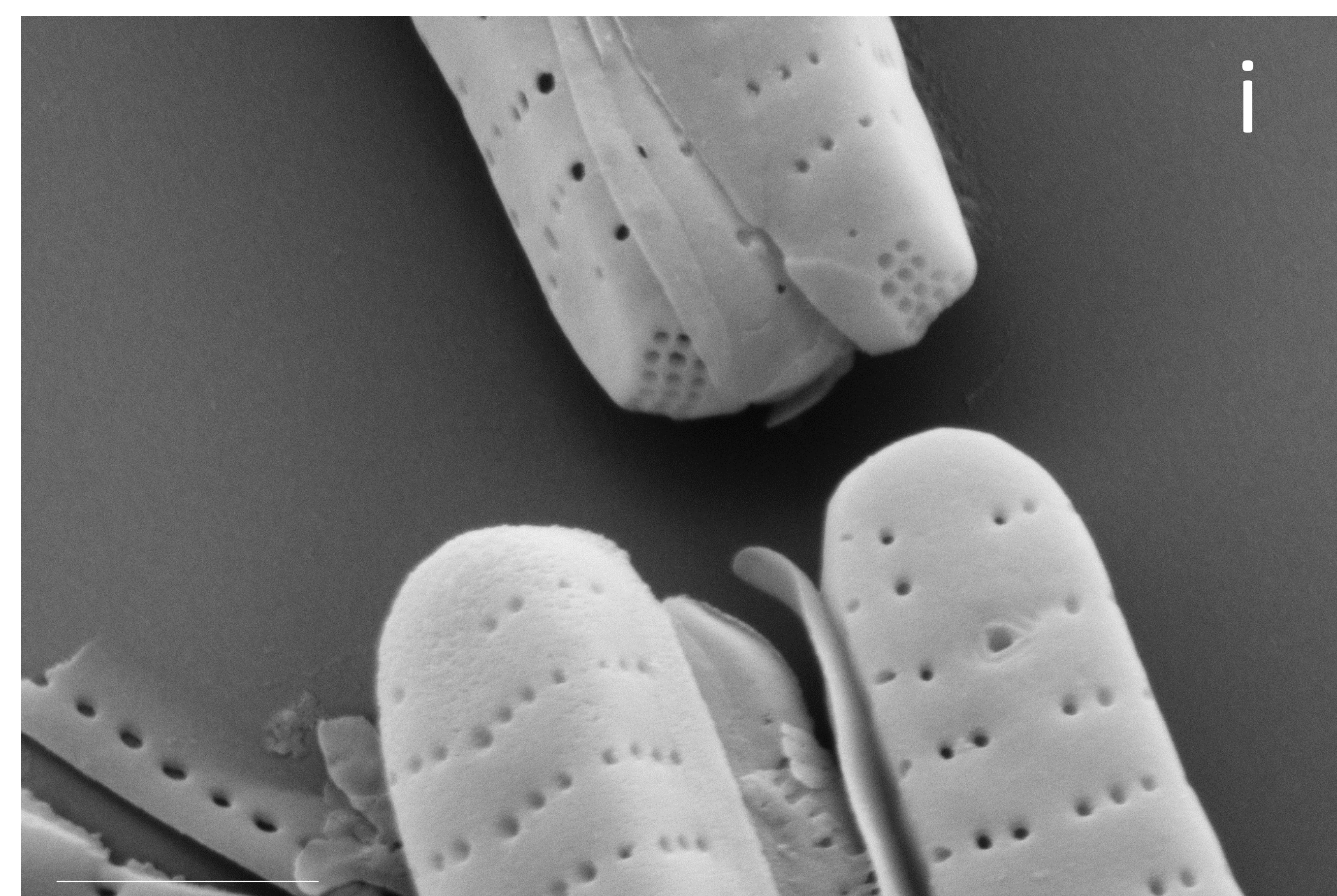
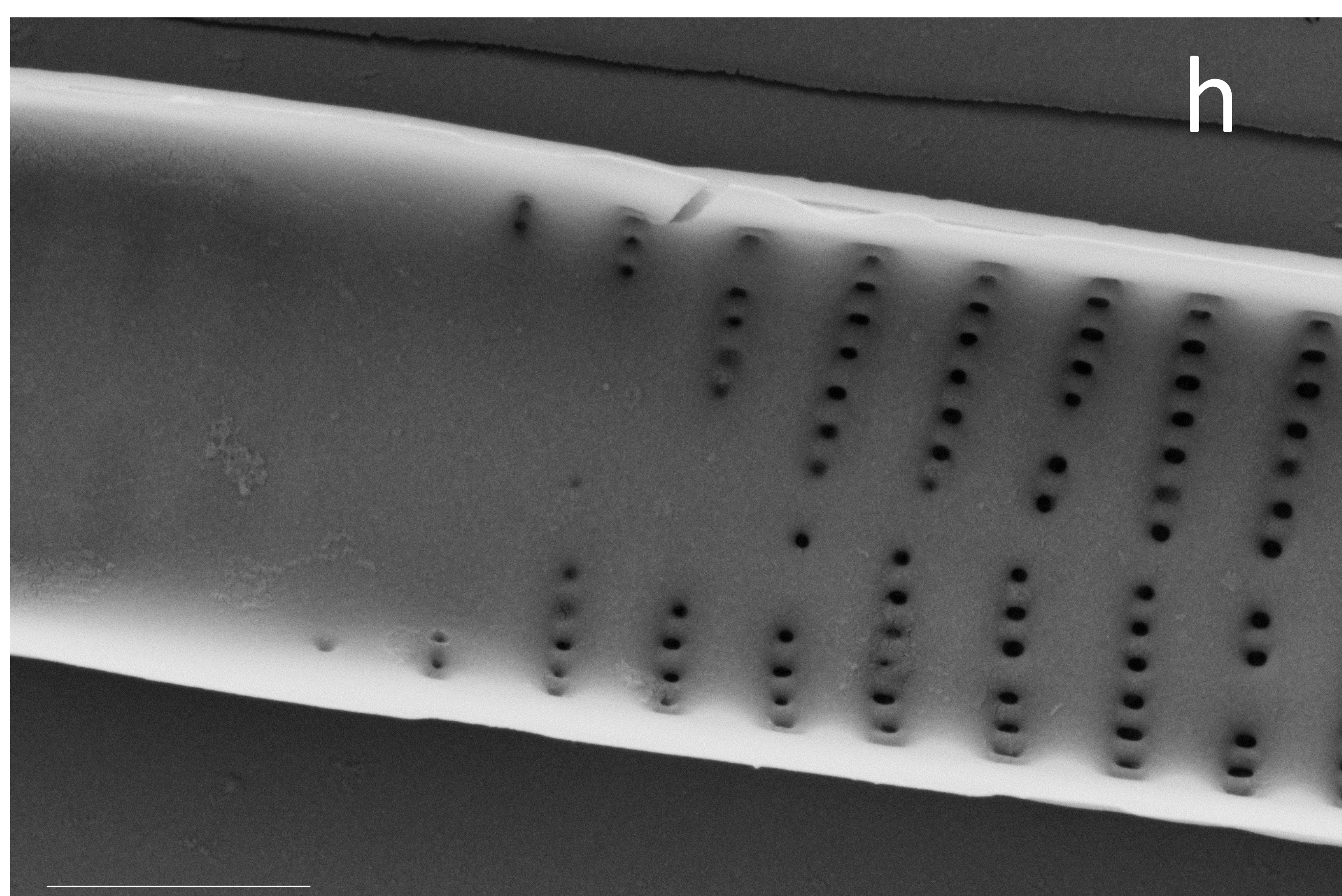
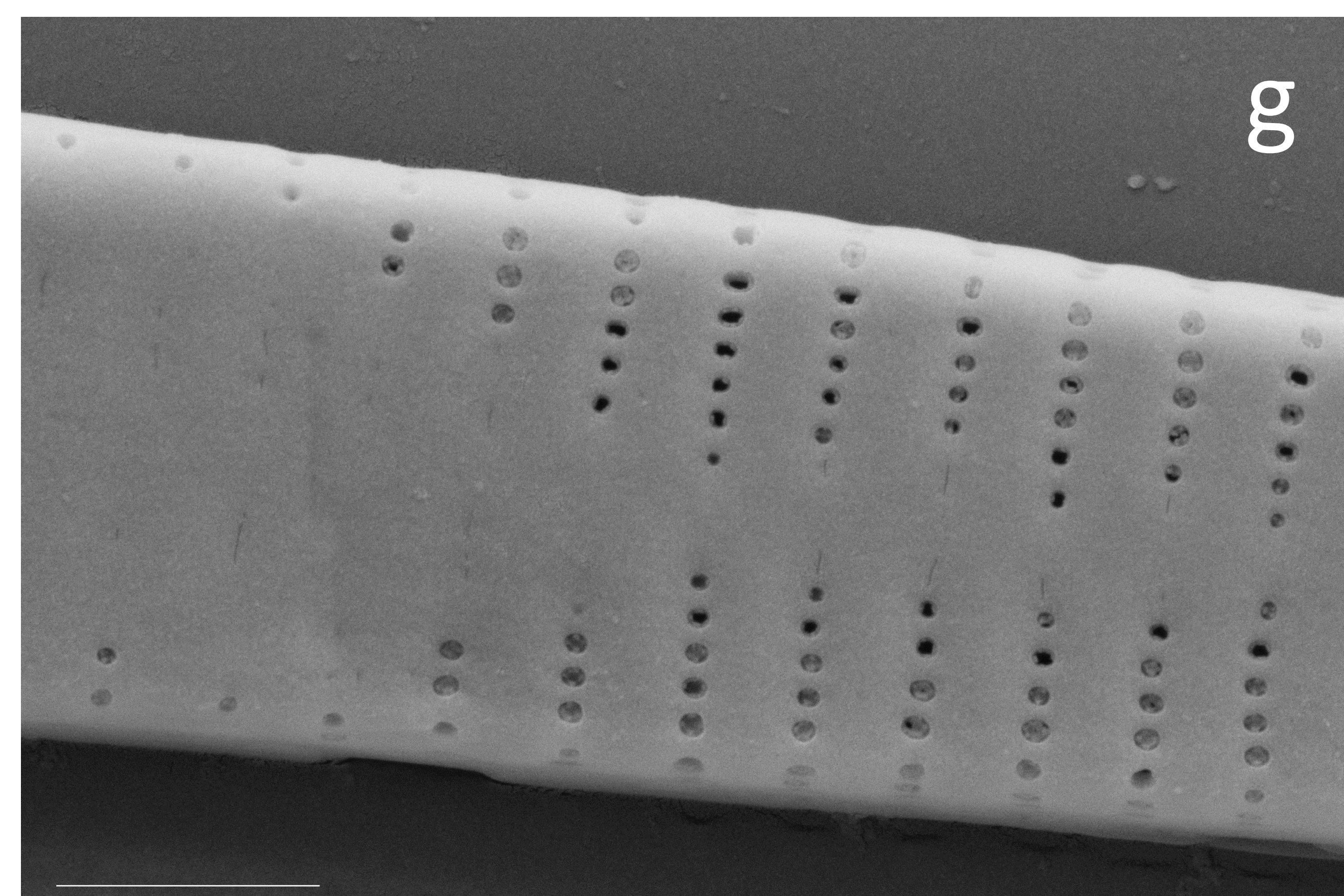
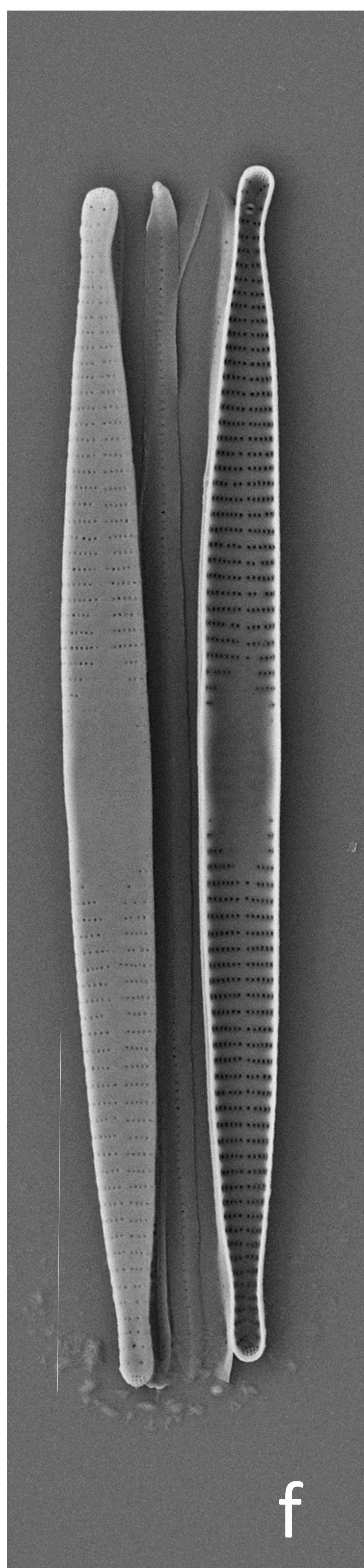
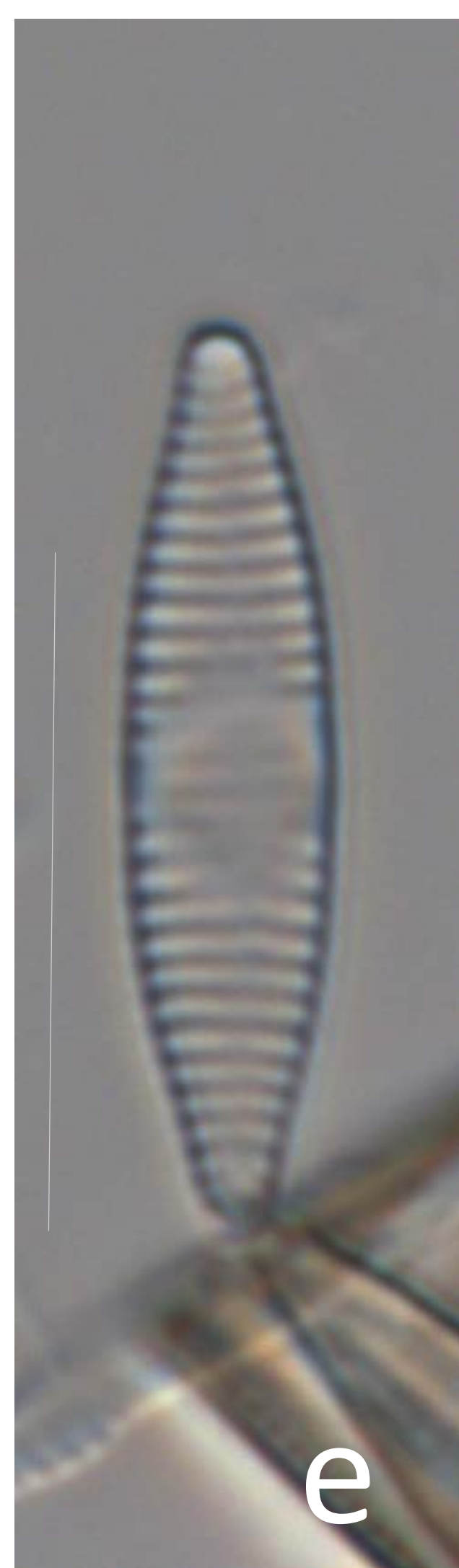
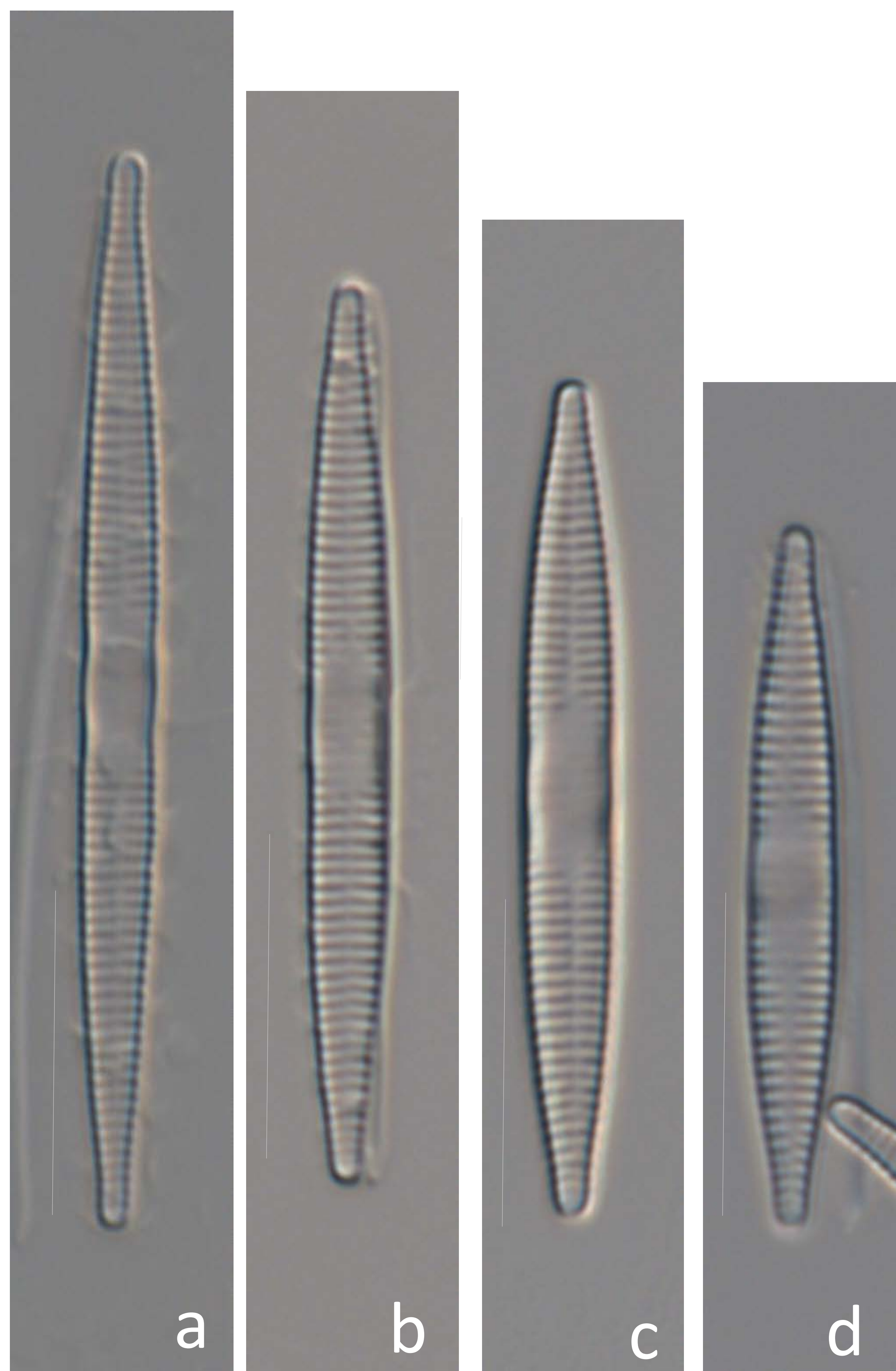
Table 1

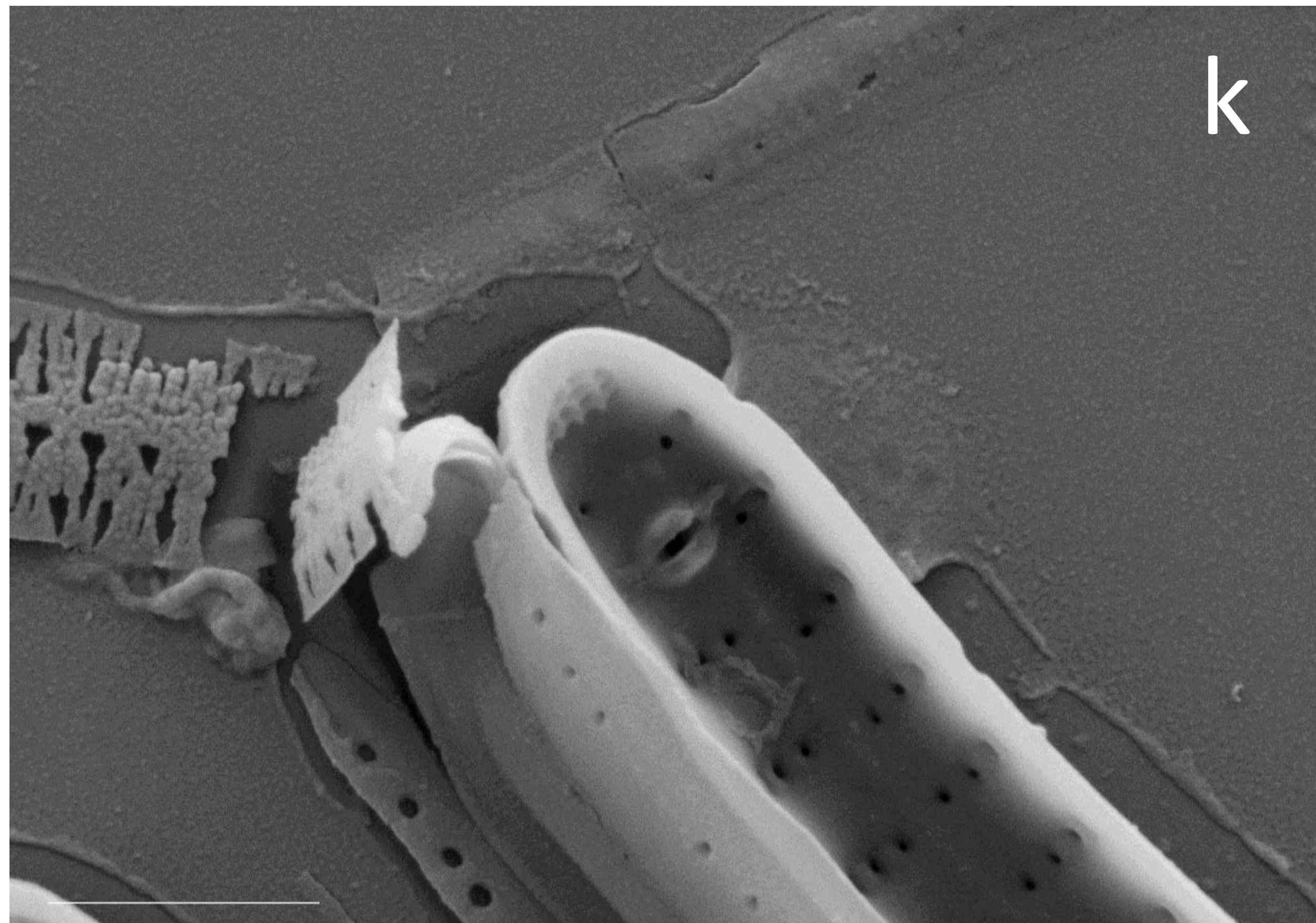
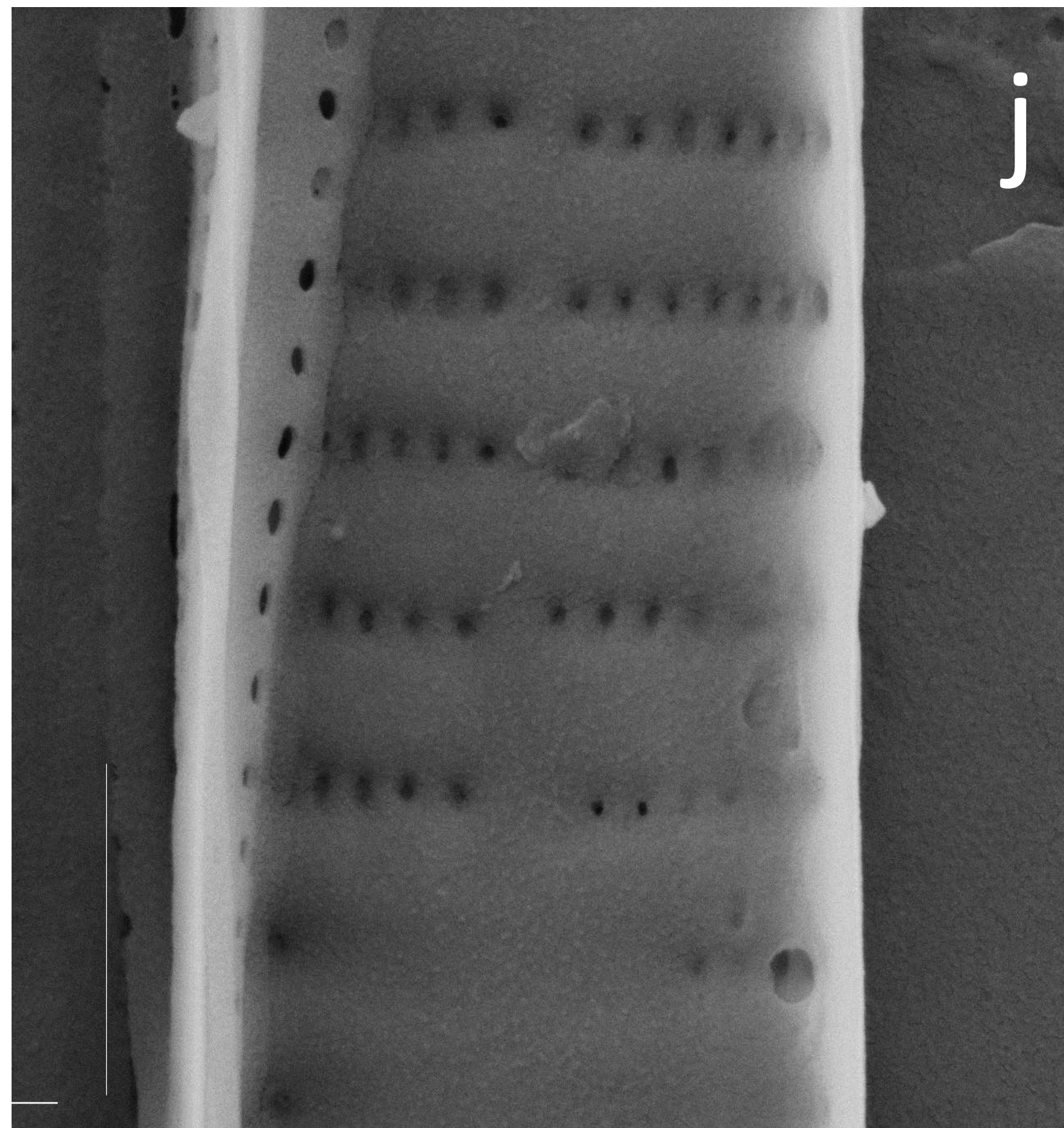
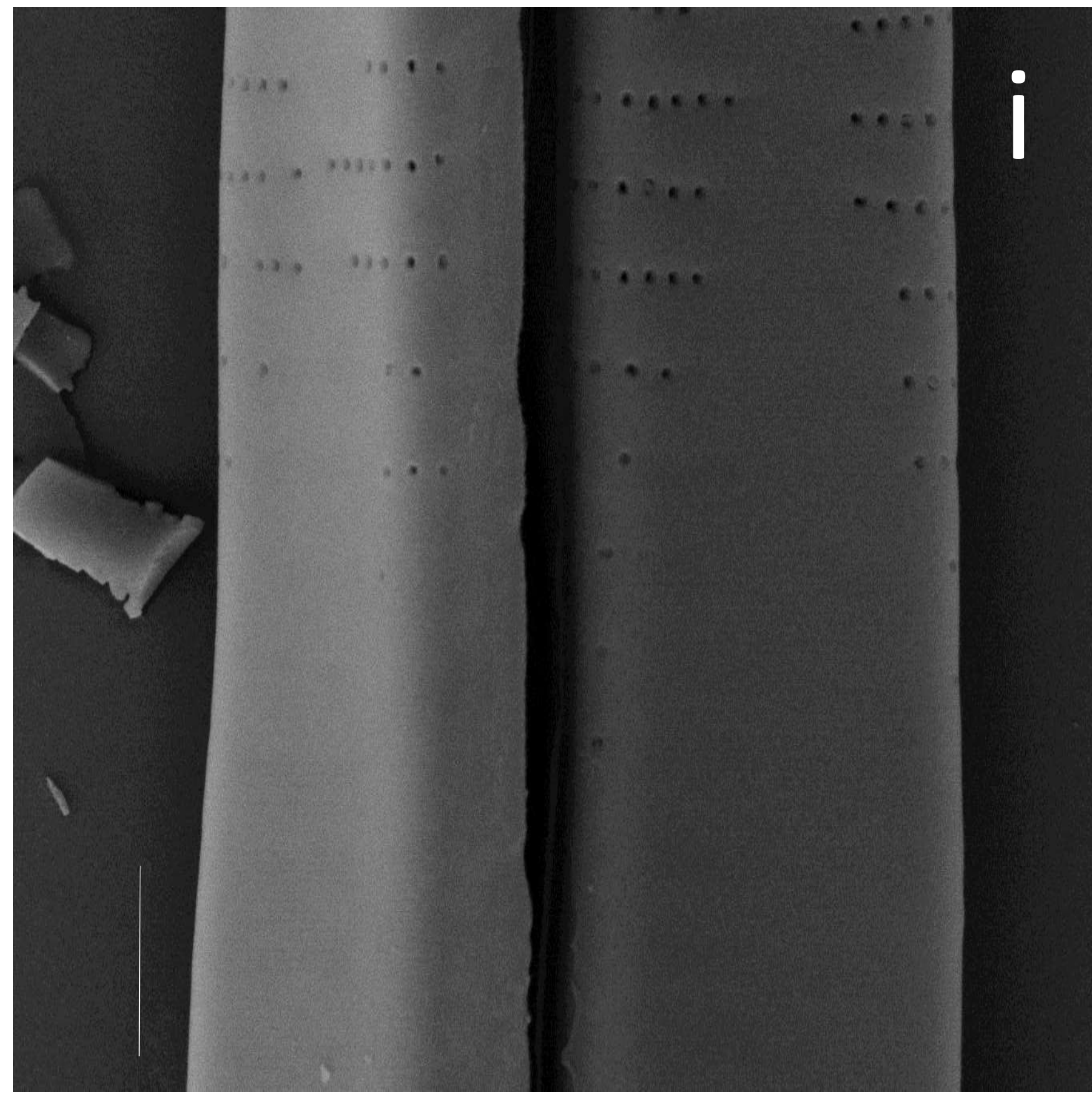
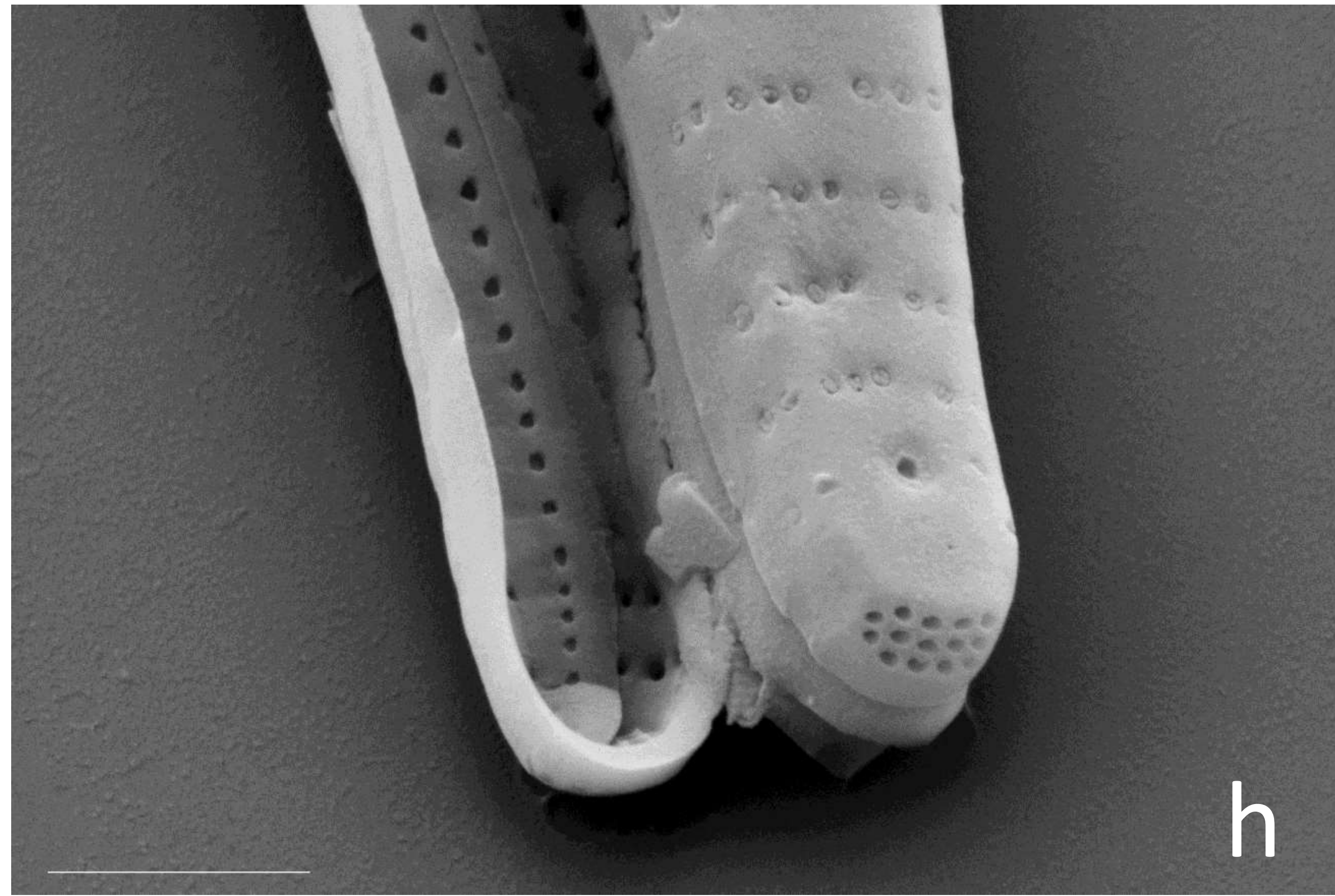
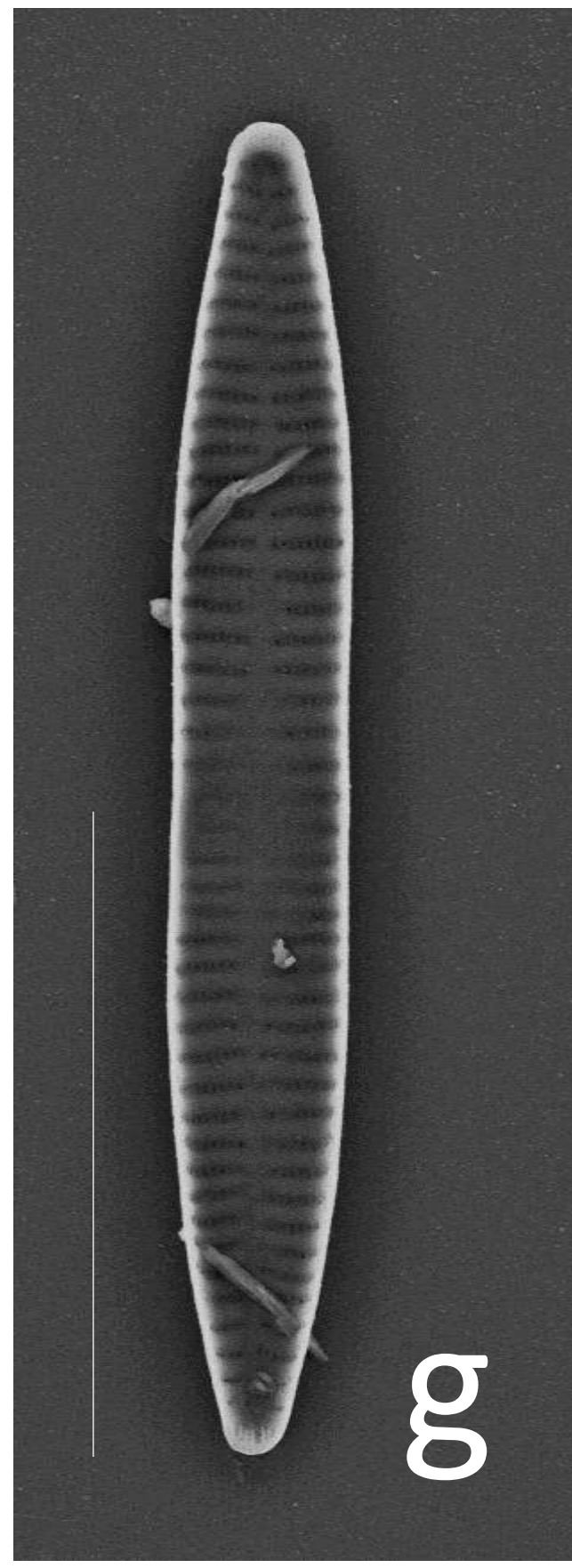
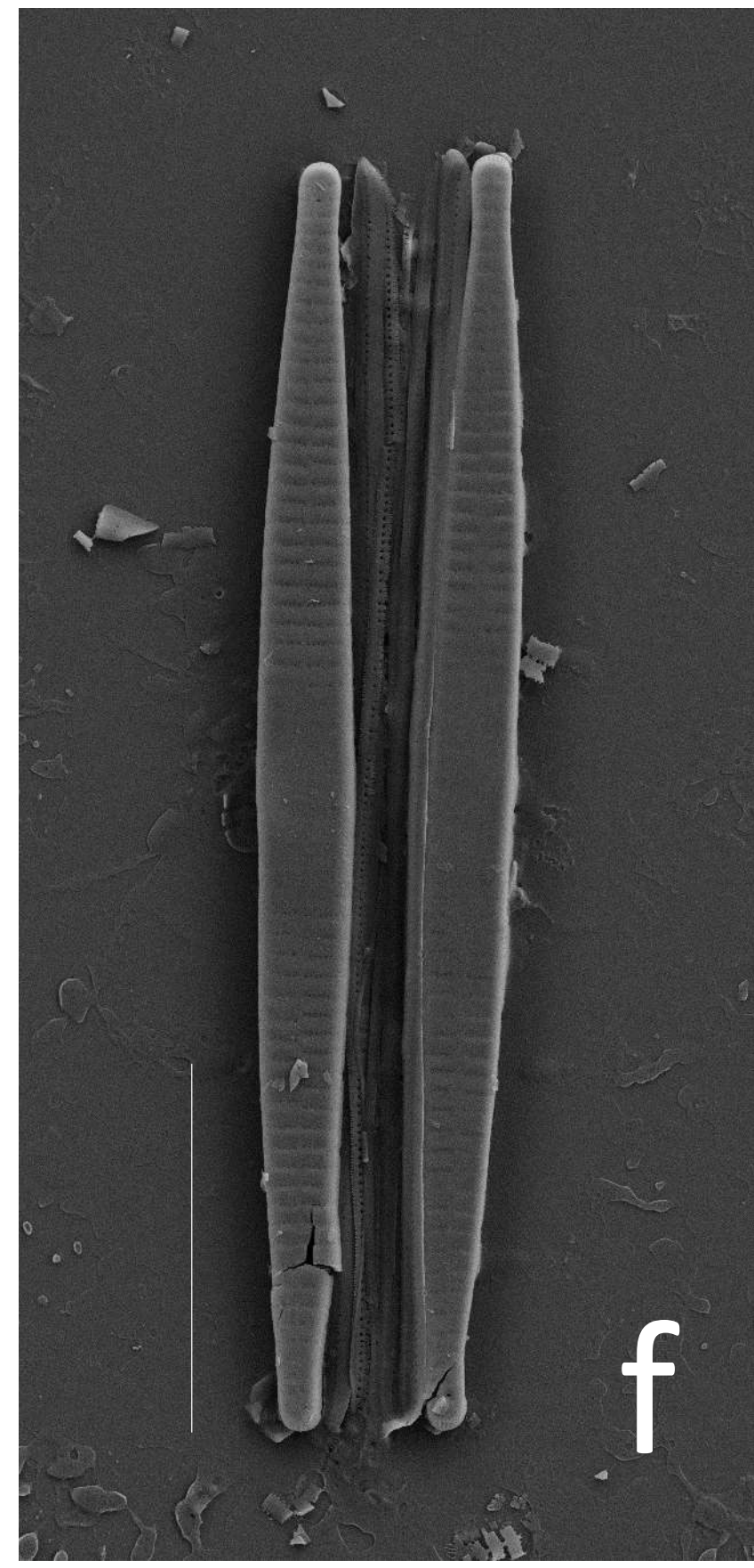
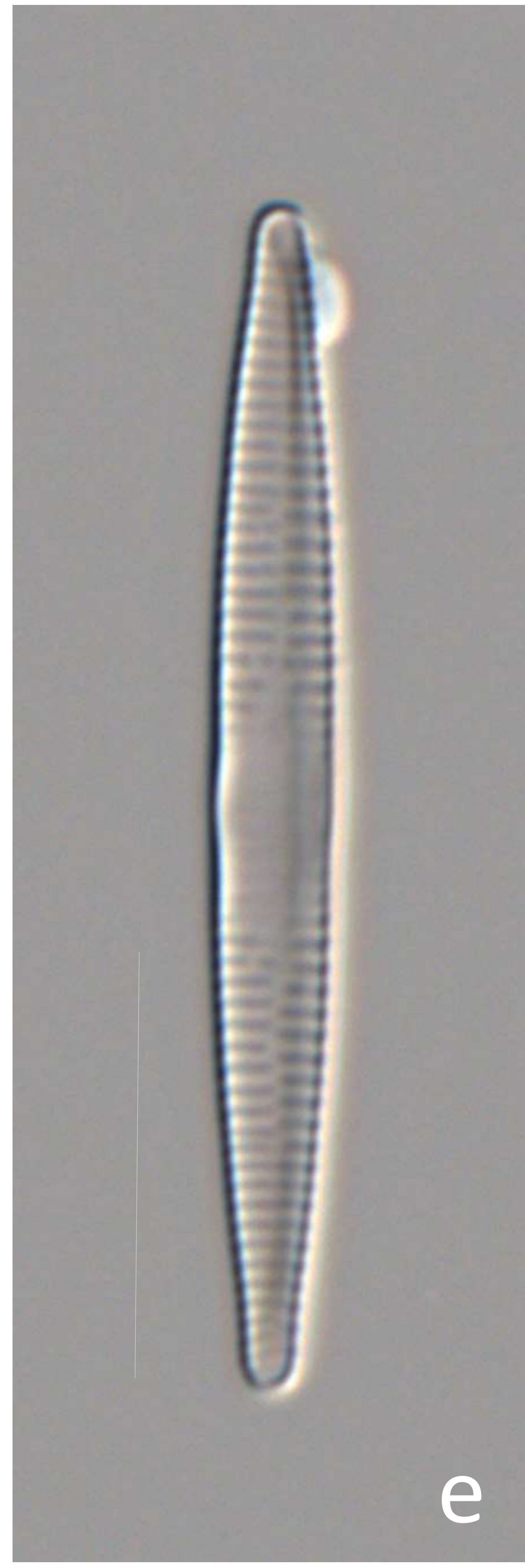
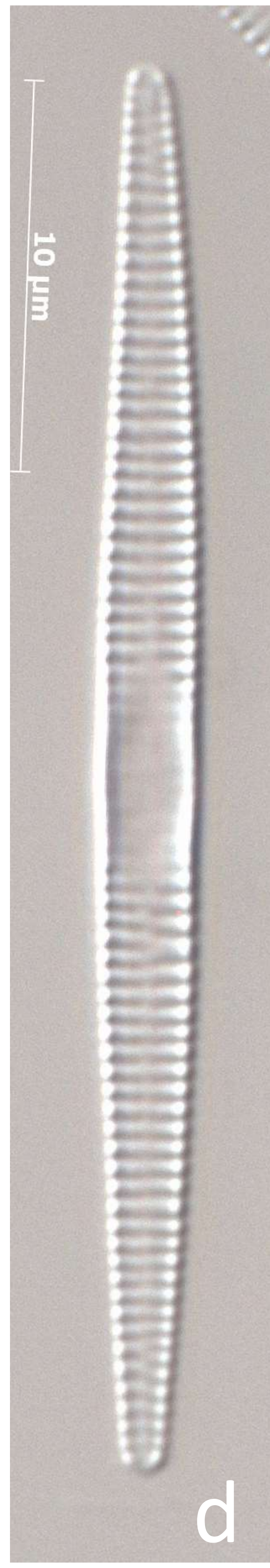
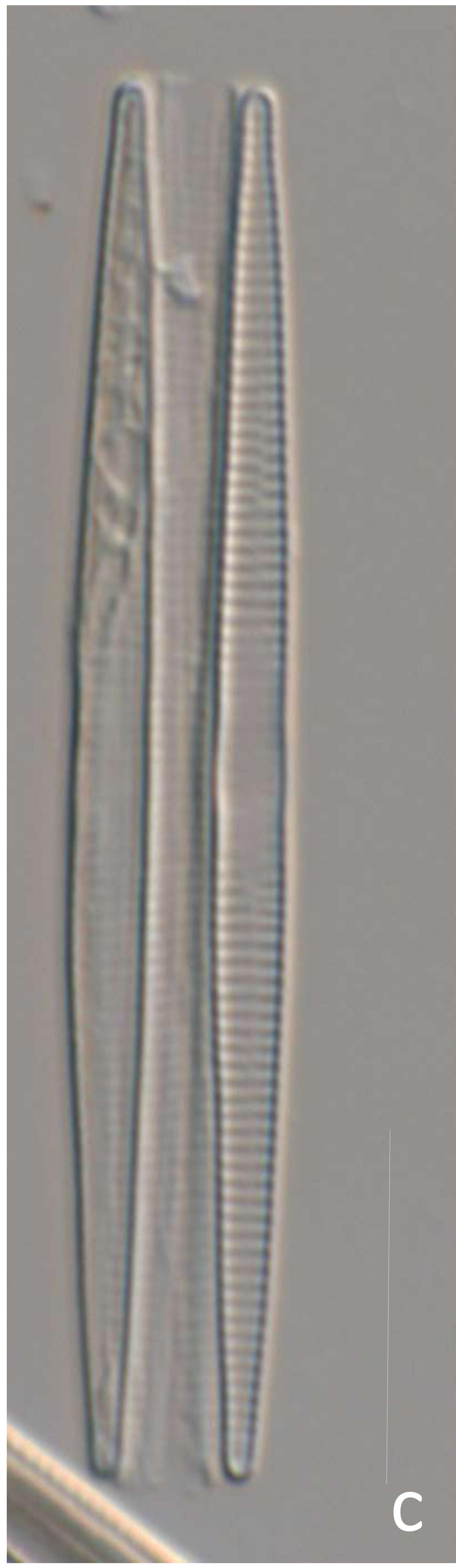
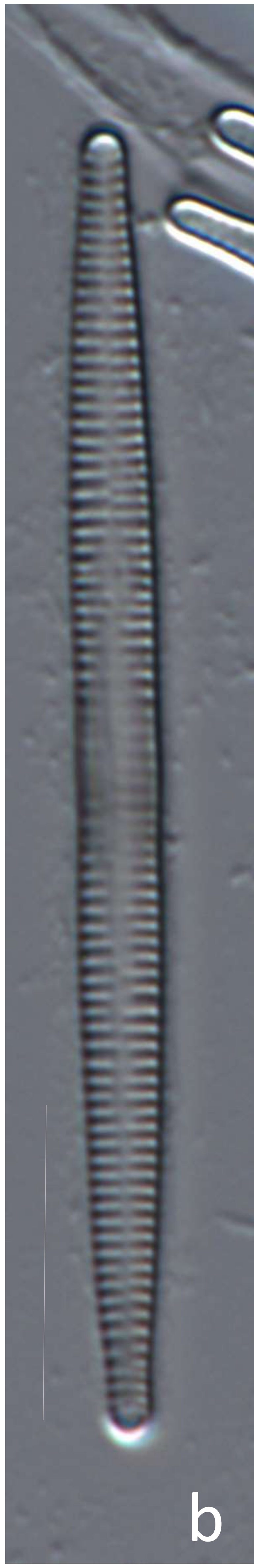
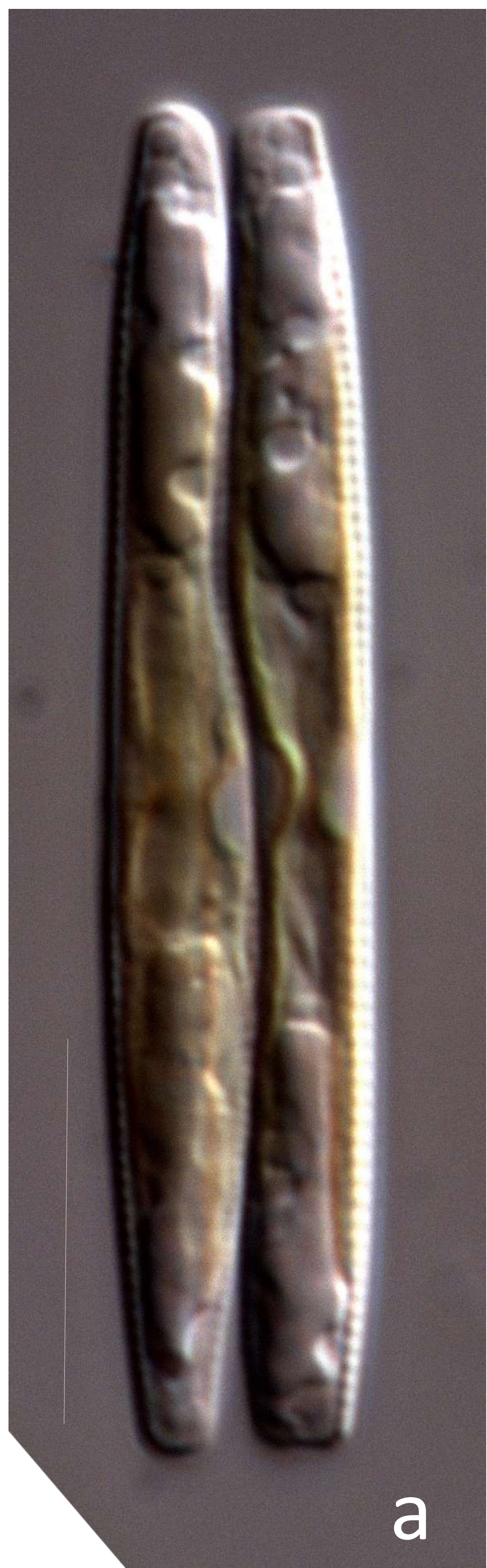
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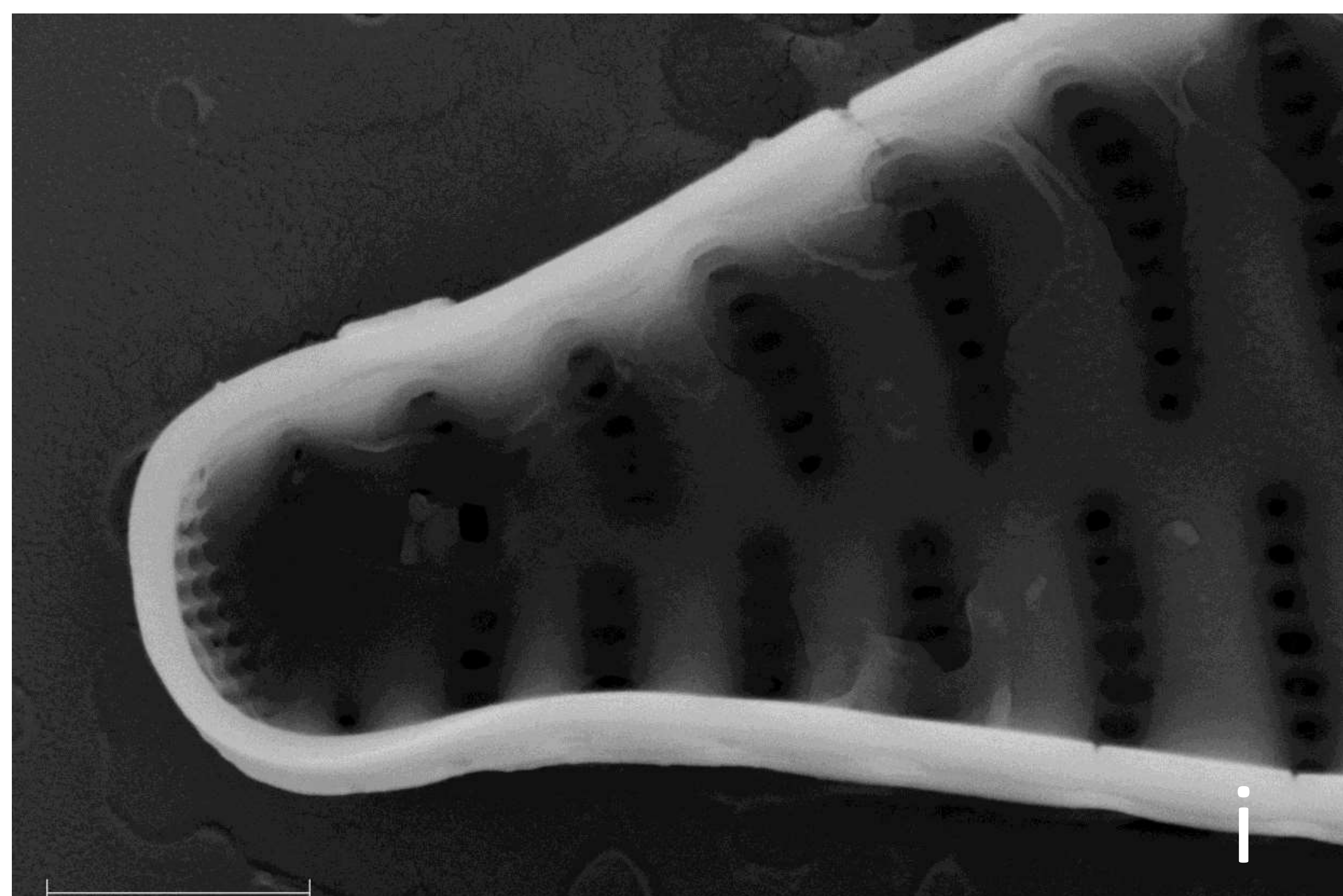
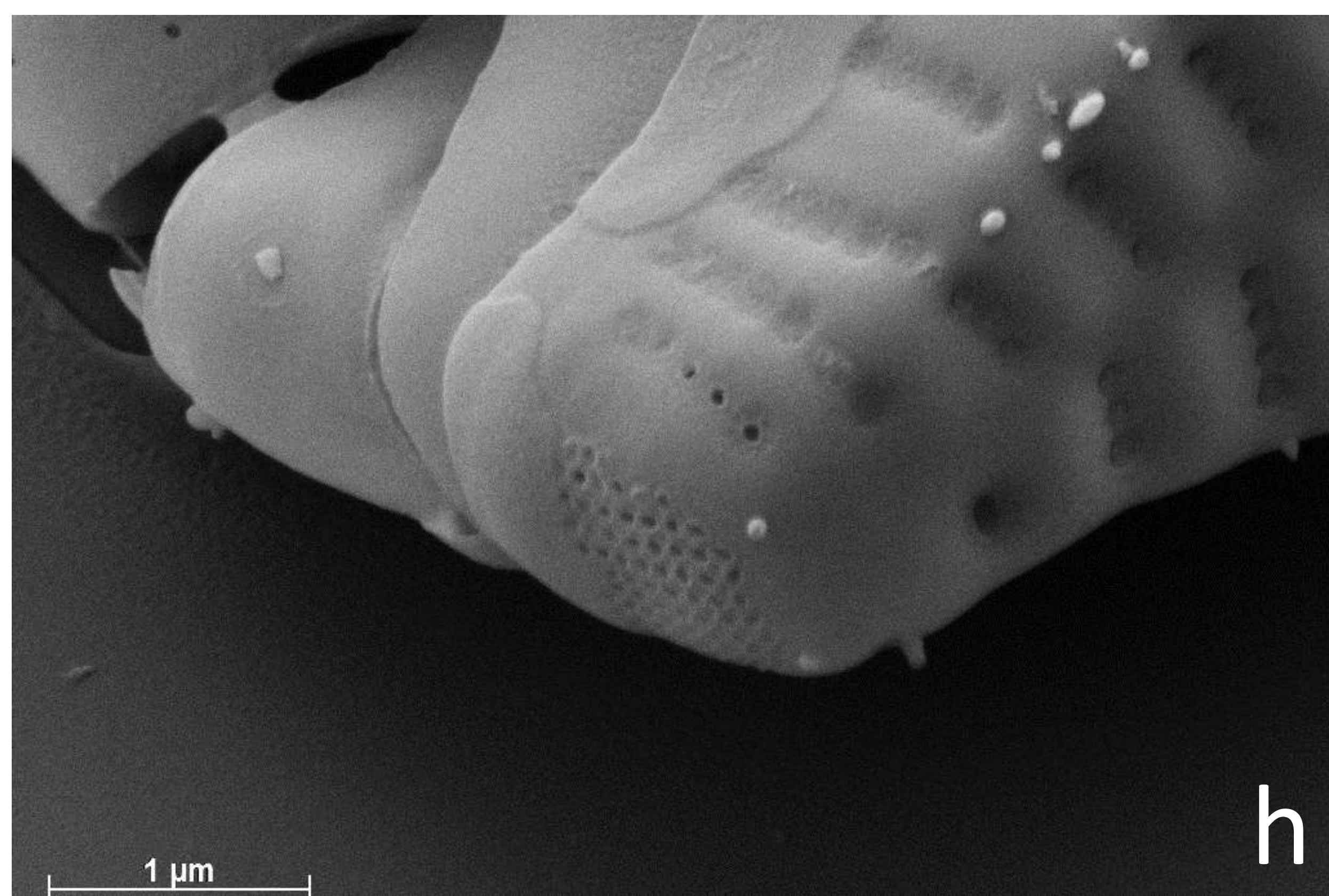
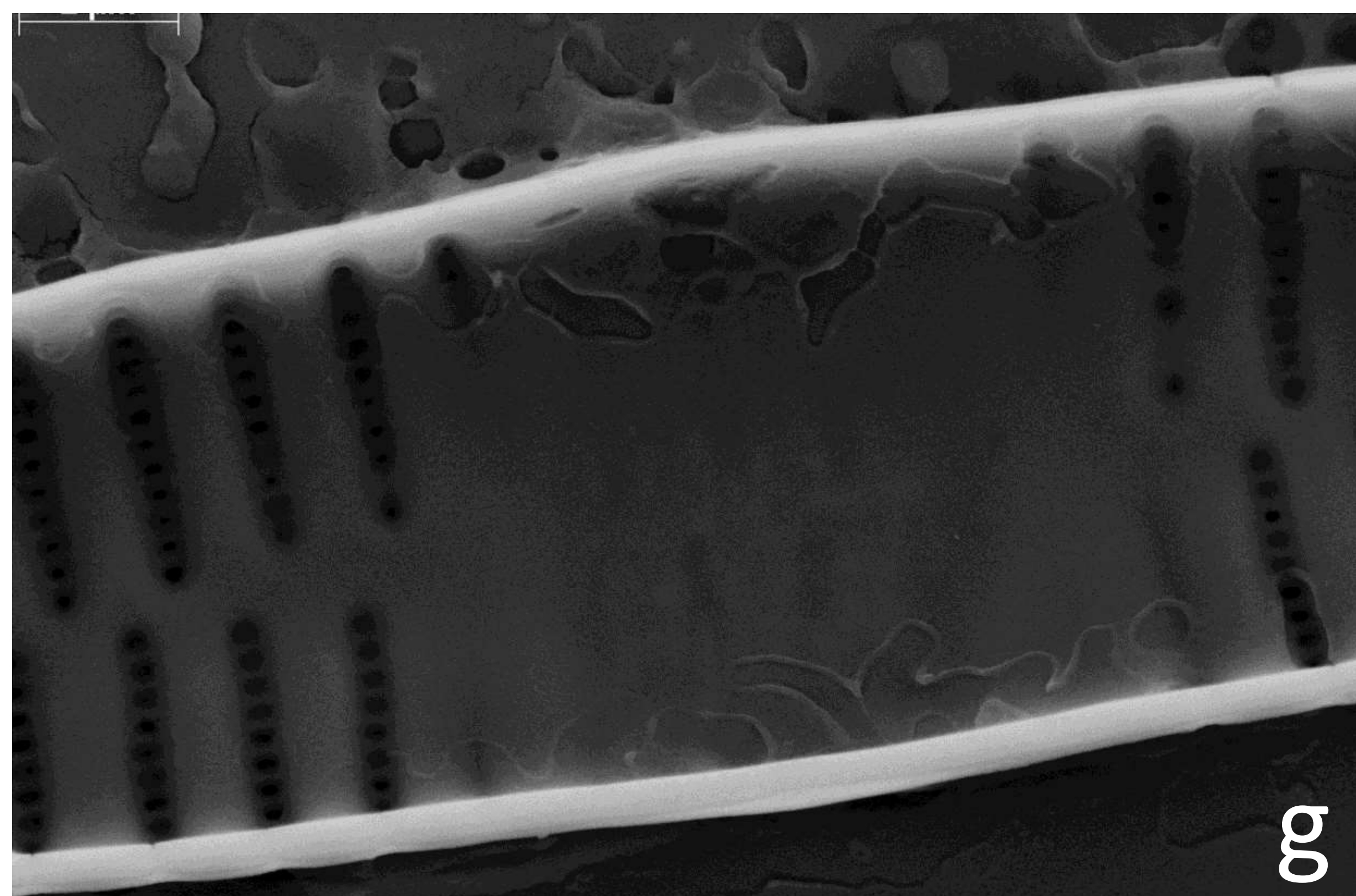
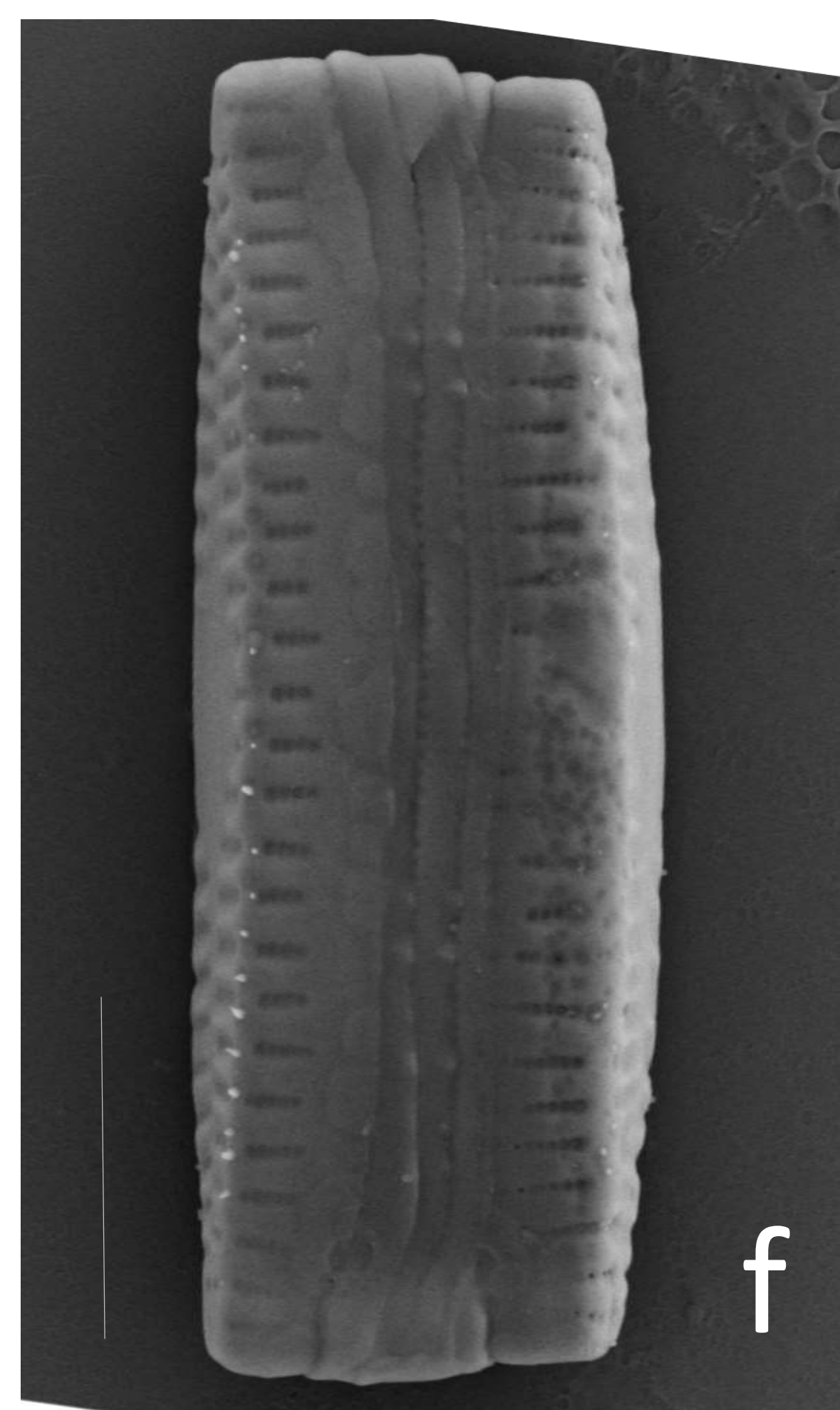
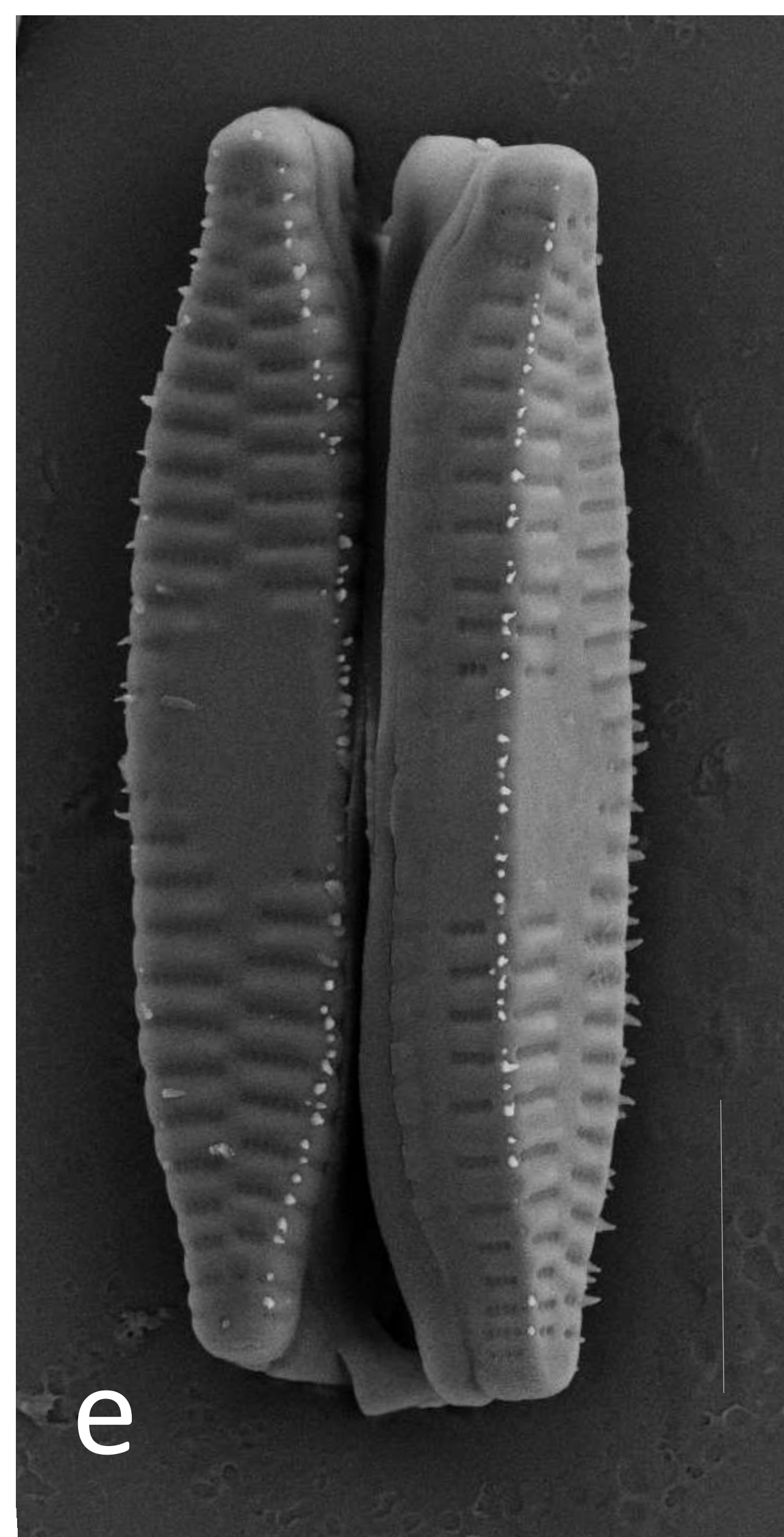
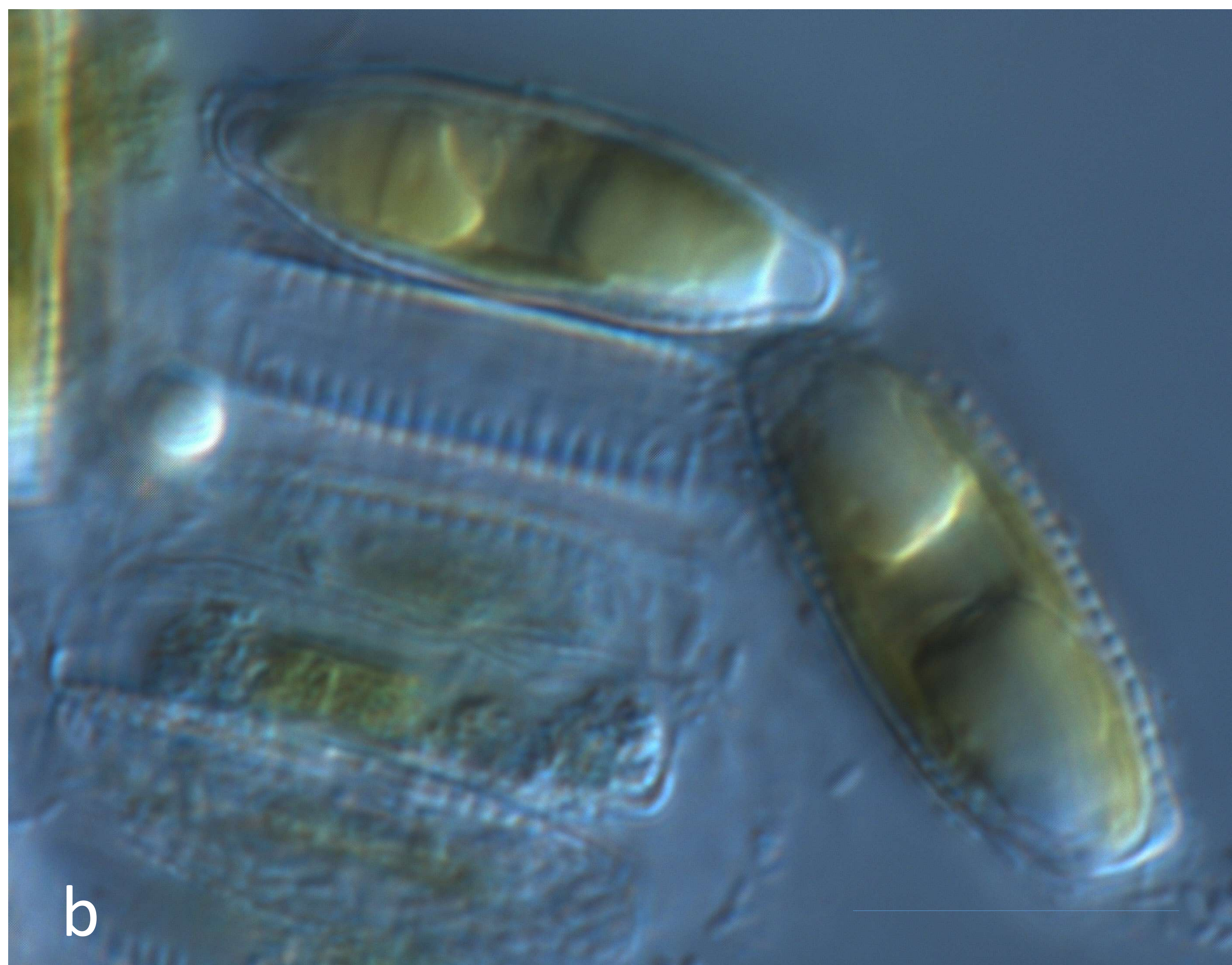
Table 2

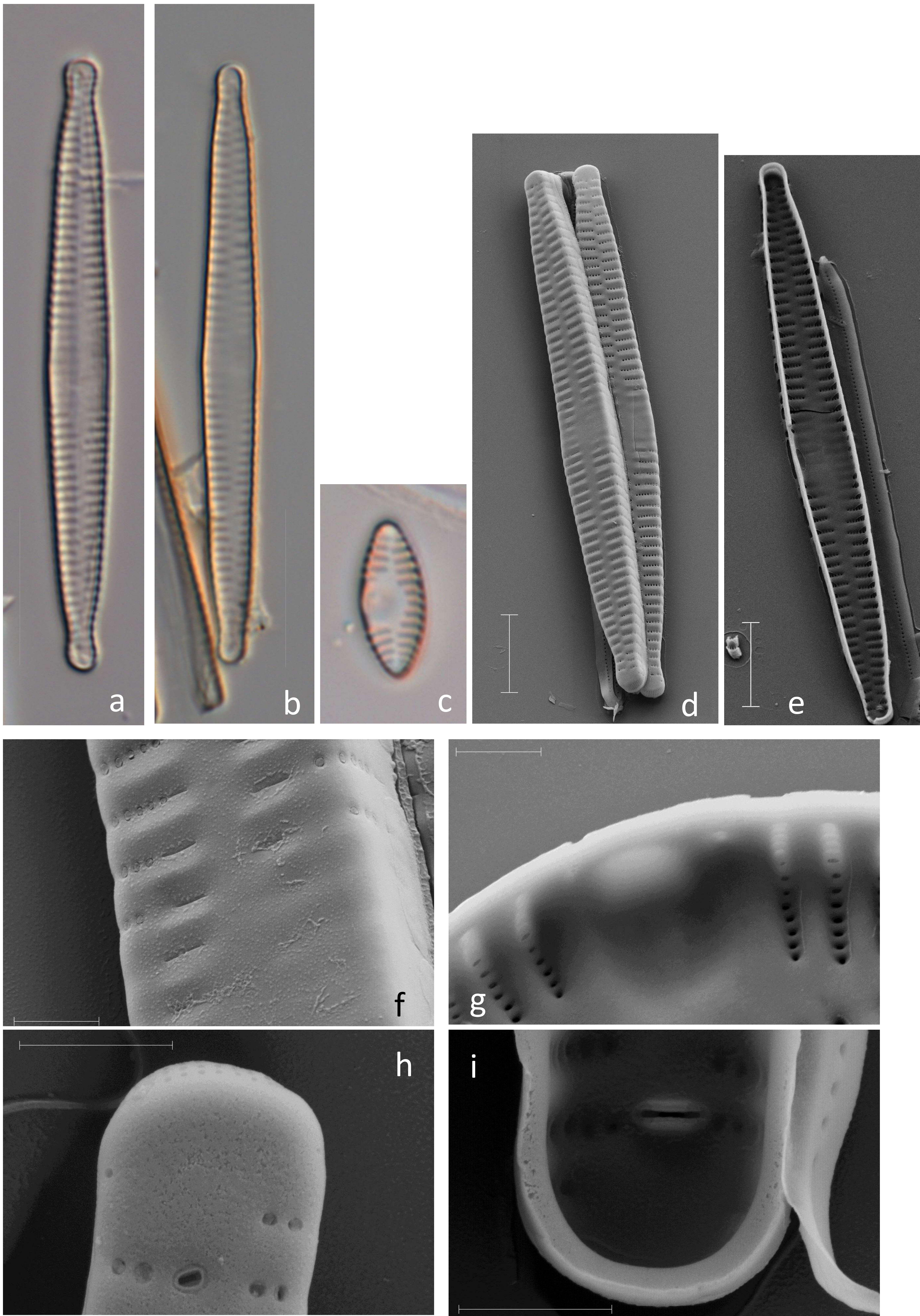
	<i>Fragilaria gracilis</i> Østrup	<i>Fragilaria gracilis</i> Østrup	<i>Fragilaria agnesiae</i> nov. spec.	<i>Fragilaria heatherae</i> nov. spec.	<i>Fragilaria joachimii</i> nov. spec.	<i>Fragilaria perminuta</i> (Grunow) Lange-Bertalot by Tuji & Williams	<i>Fragilaria subconstricta</i> Østrup	<i>Fragilaria tenera</i> (W. Smith) Lange-Bertalot	<i>Fragilaria</i> "sp. 1"
working name in ms	FRGA2	FRGA1	FVAU	FCAP1	FCAP2	FPEM	FTNS	FTEN1	FTEN2
nr of strains studied for SEM	9	10	1	3	7	5	2	6	2
Rimoportul	number of rimoportulae	1	1	1	1	1 (2 in one specimen)	1	1	1
	position of rimoportulae	on valve face at one of the poles	on valve face at one of the poles	on valve face at one of the poles	on valve face at one of the poles	on valve face at one of the poles close to final pore	on mantle face junction	on valve face at one of the poles close to final pore	on valve face at one of the poles close to final pore
Apical pore fields	number of apical pore fields	2	2	2	2	2	2	2	2
	form of apical pore fields	oval to rectangular	oval to rectangular	rectangular	rectangular	rectangular	oval to rectangular	rectangular	triangular to rectangular
	number of columns per field	4-8	4-8	8-10	long strains: 12-14; short strain: 9	5-14	6-10	15	5-7
	maximum number of pores per column	4	4	5	6	5	6	5	4
	Spines	absent	absent	irregular arranged, tiny to pyramidal with a round basis ending in a round tip	absent	irregular arranged, very tiny spines in long cells, absent in short cells	absent	gross spatulate	very regular arranged, pyramidal, often with a sharp bent tip
	Central area	rhombic	rhombic	no internal rimmed depression observed	internal rimmed depression in short cells, not observed in long cells	various grades of internal rimmed depression, clearest in short cells	clear internal rimmed depression	no observations	rhombic with external ghost striae
molecular phylogeny	cluster support [%]	97% (as subgroup within FGRA)	69% (FGRA1 & FGRA2 together)	98%	79%	84%	87%	79%	100%
	Evolutionary Divergence between Sequences (direct MEGA output)	0-0.001	0-0.004	0	0-0.003	0-0.002	0-0.002	0-0.001	0-0.001
	nr of haplotypes	2	5	1	2	2	3	2	2



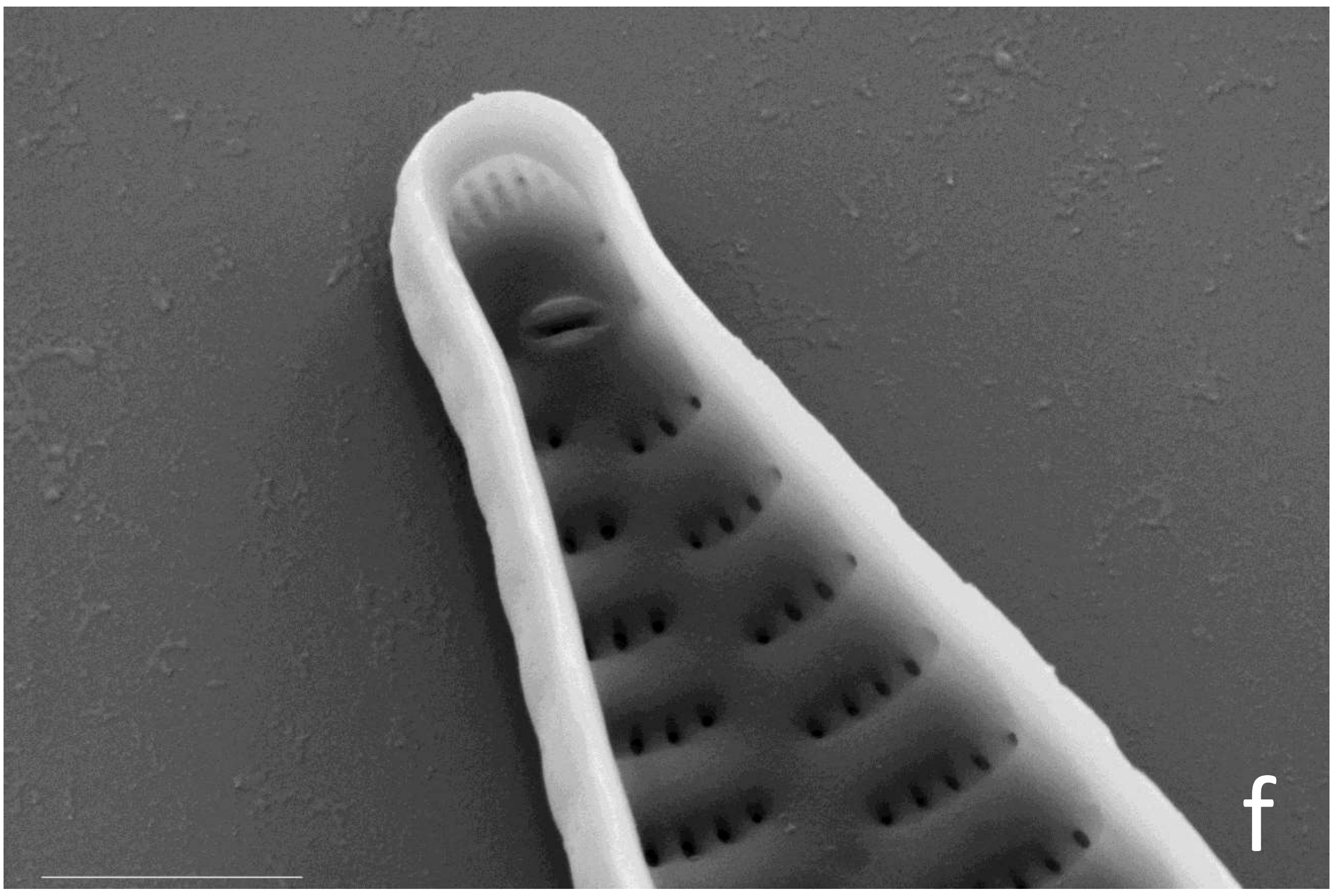
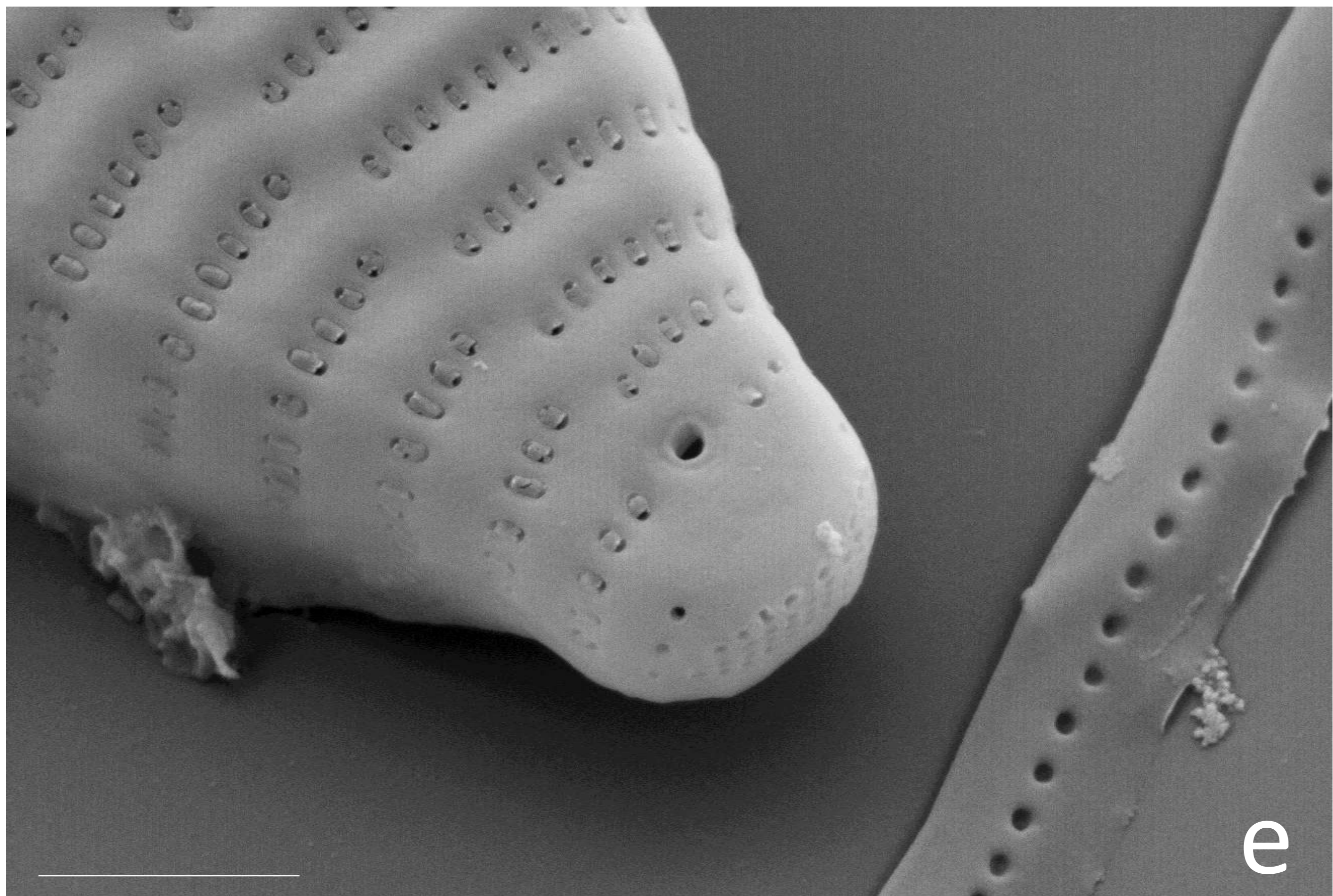
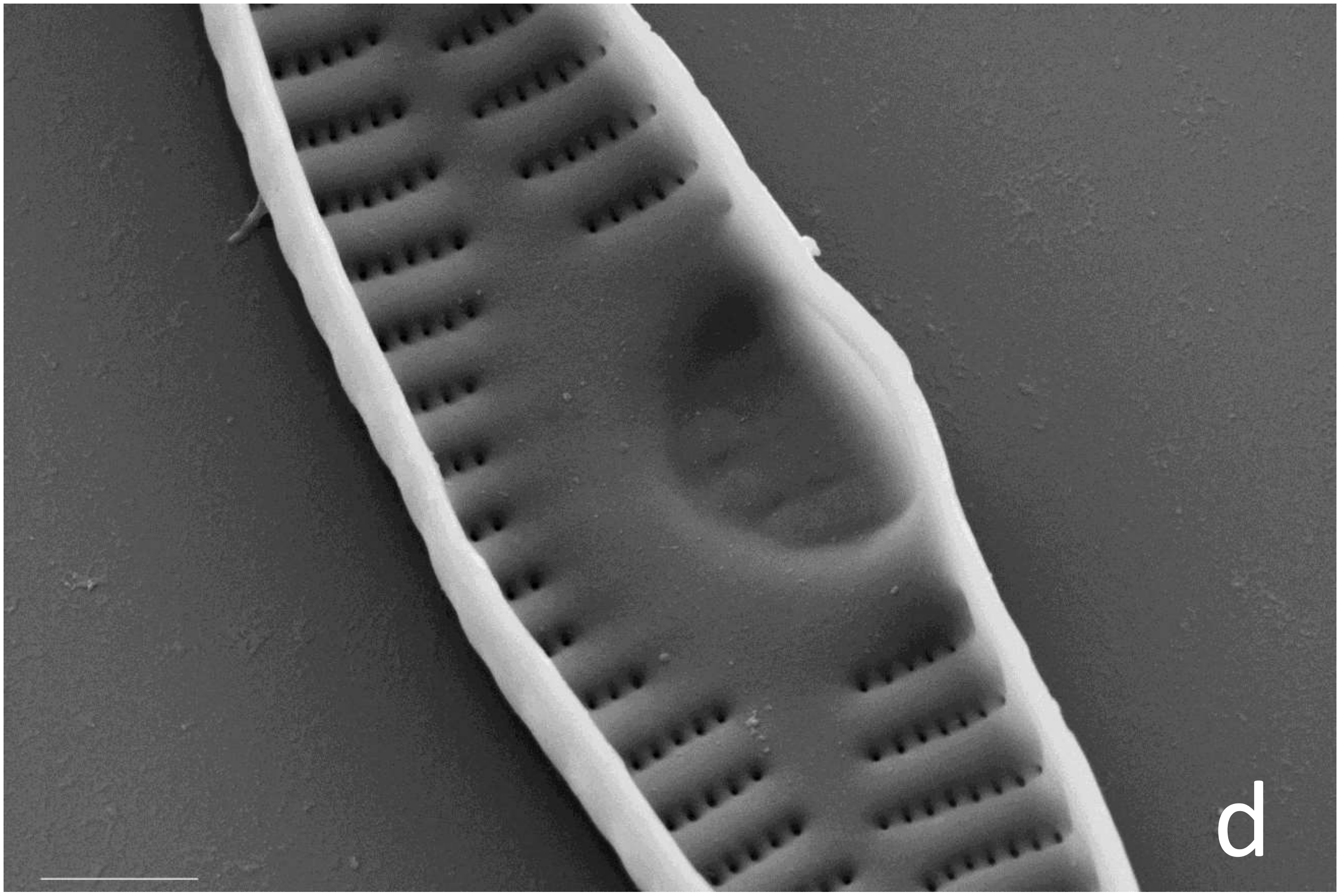
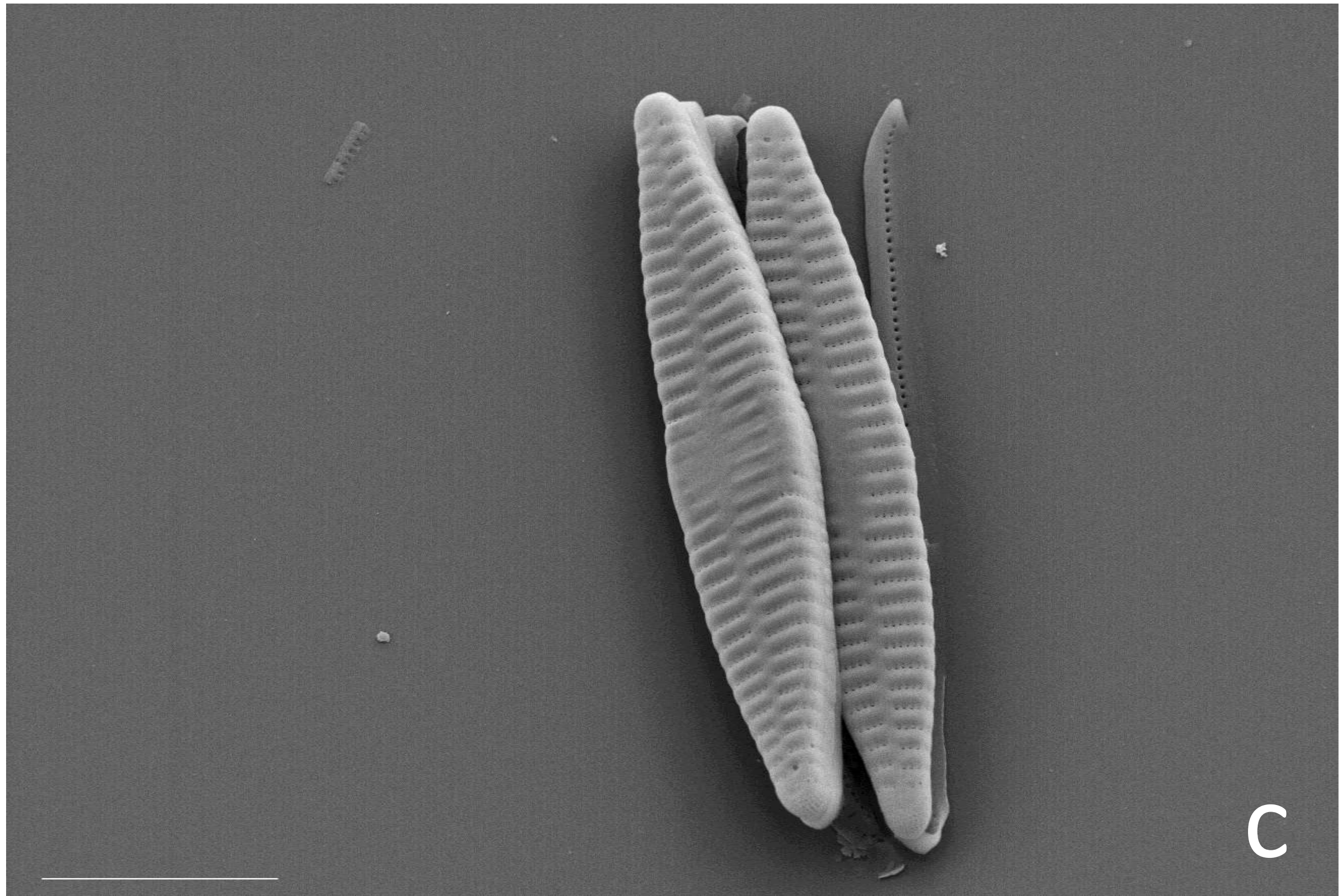
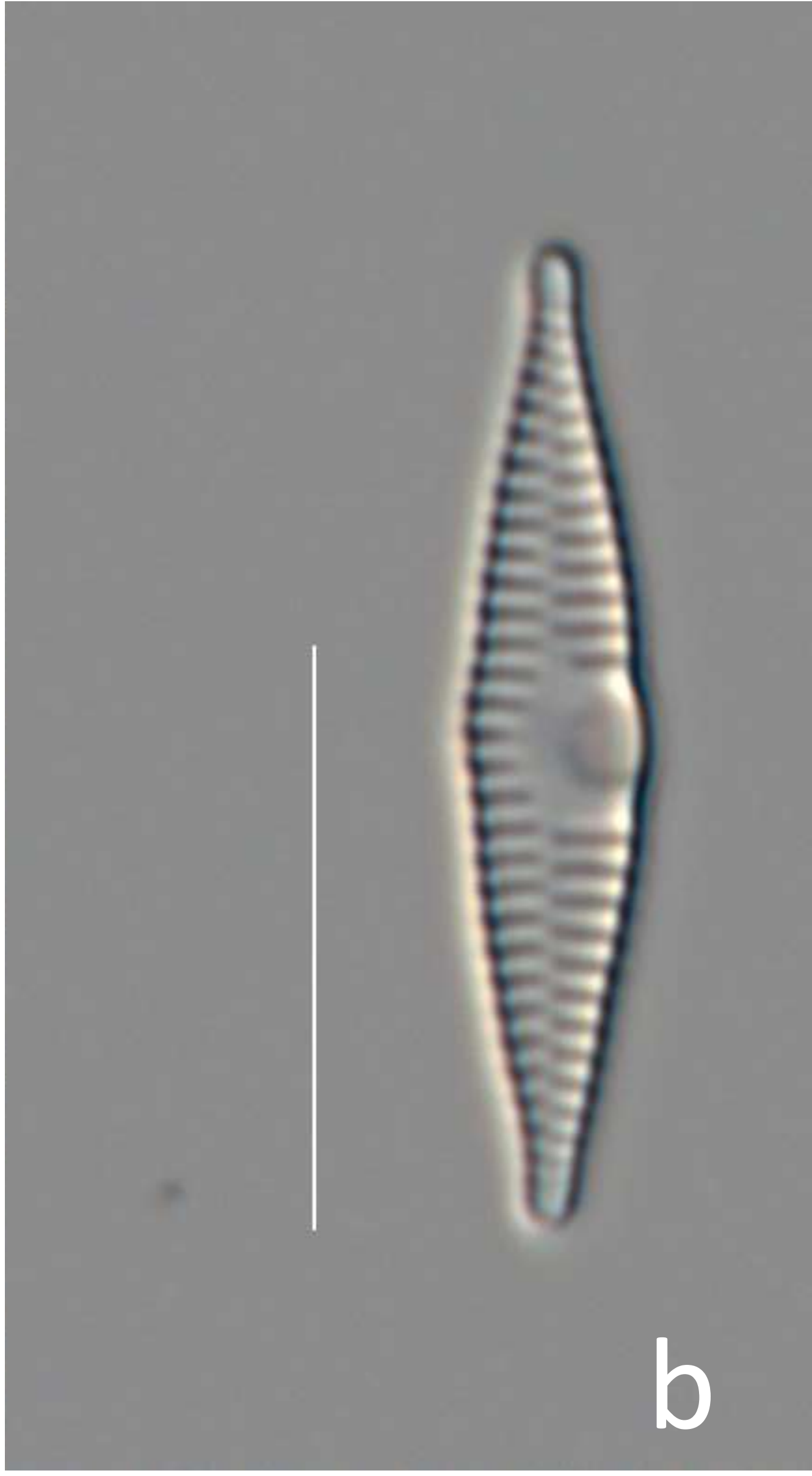
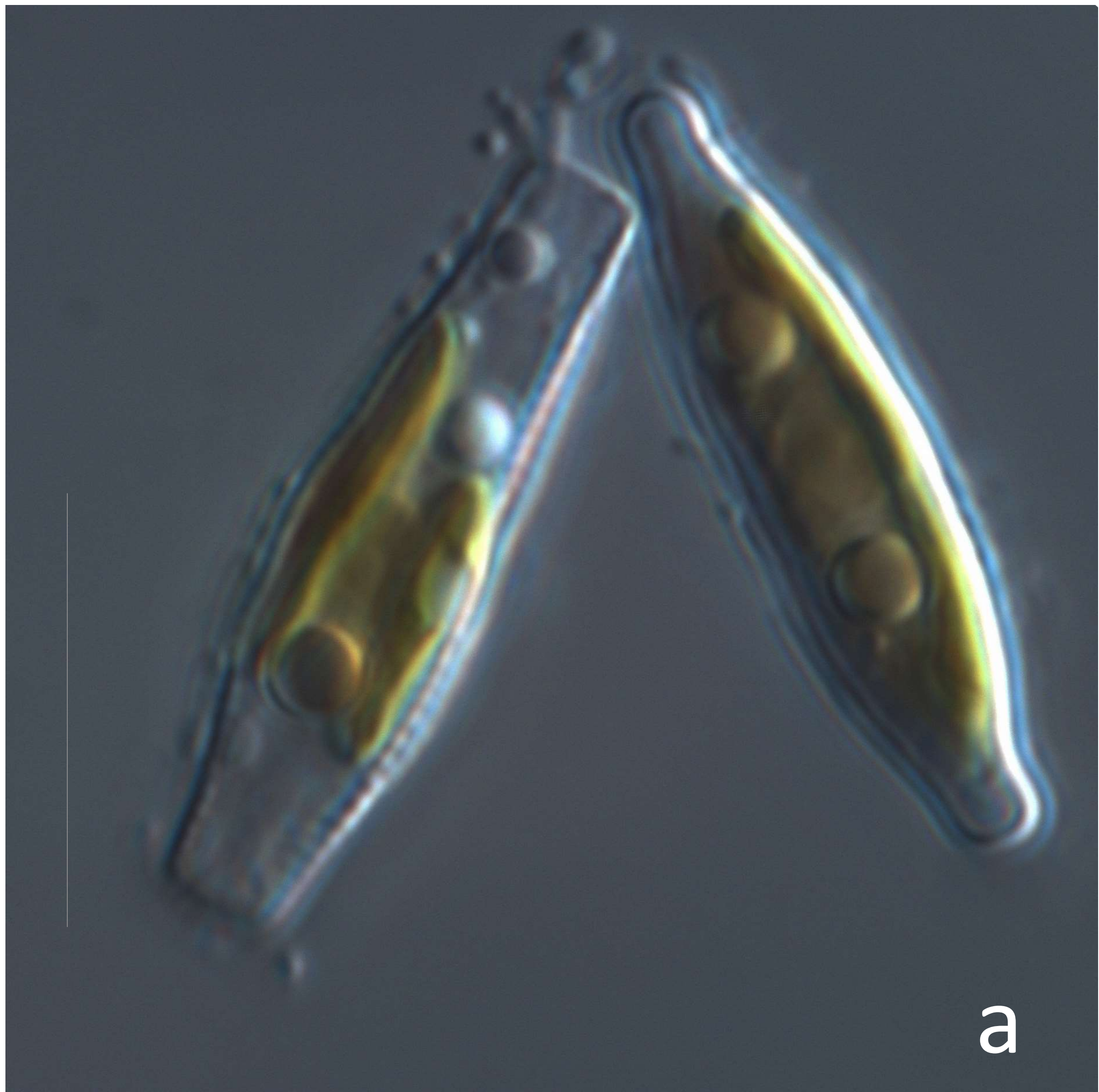


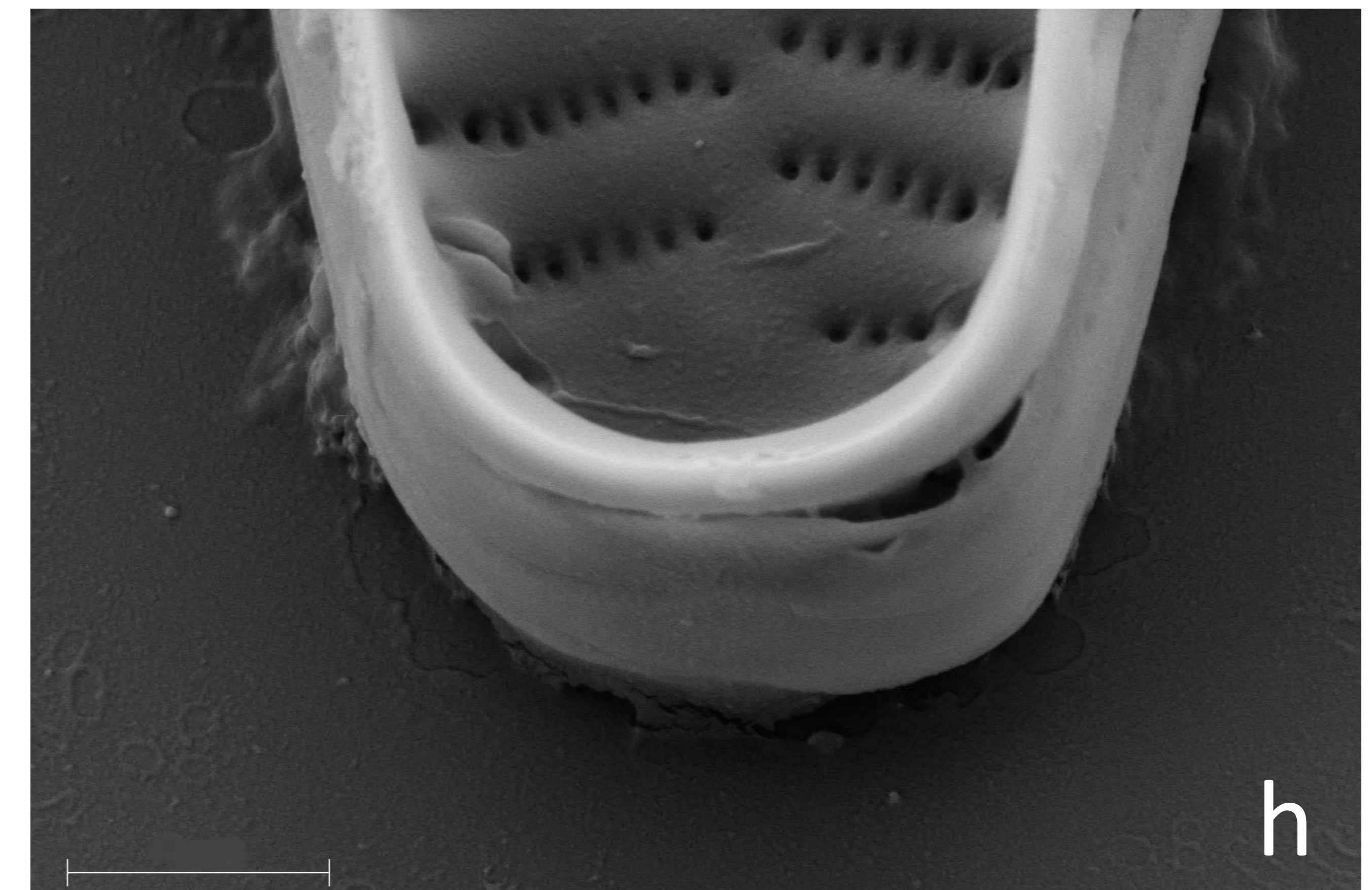
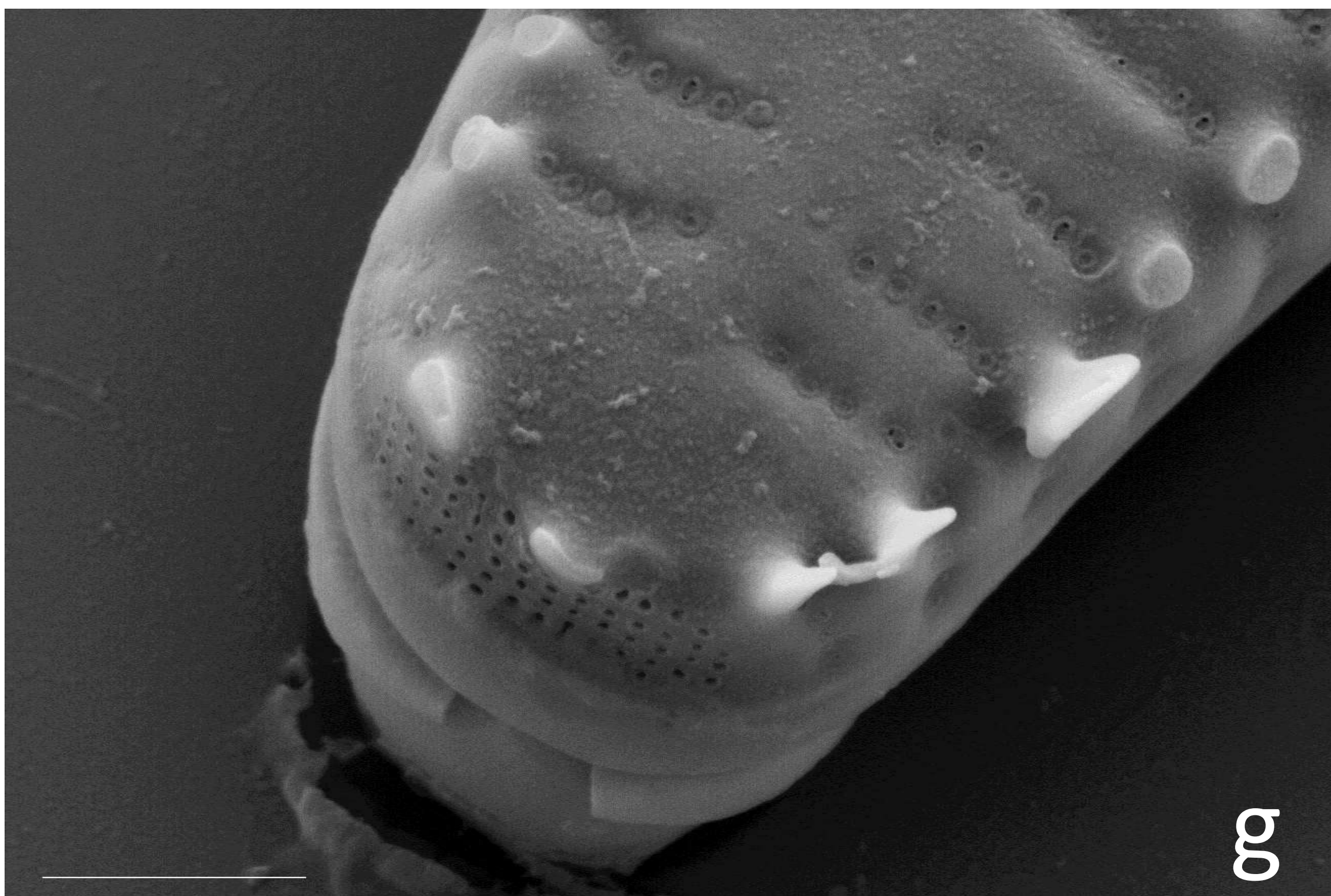
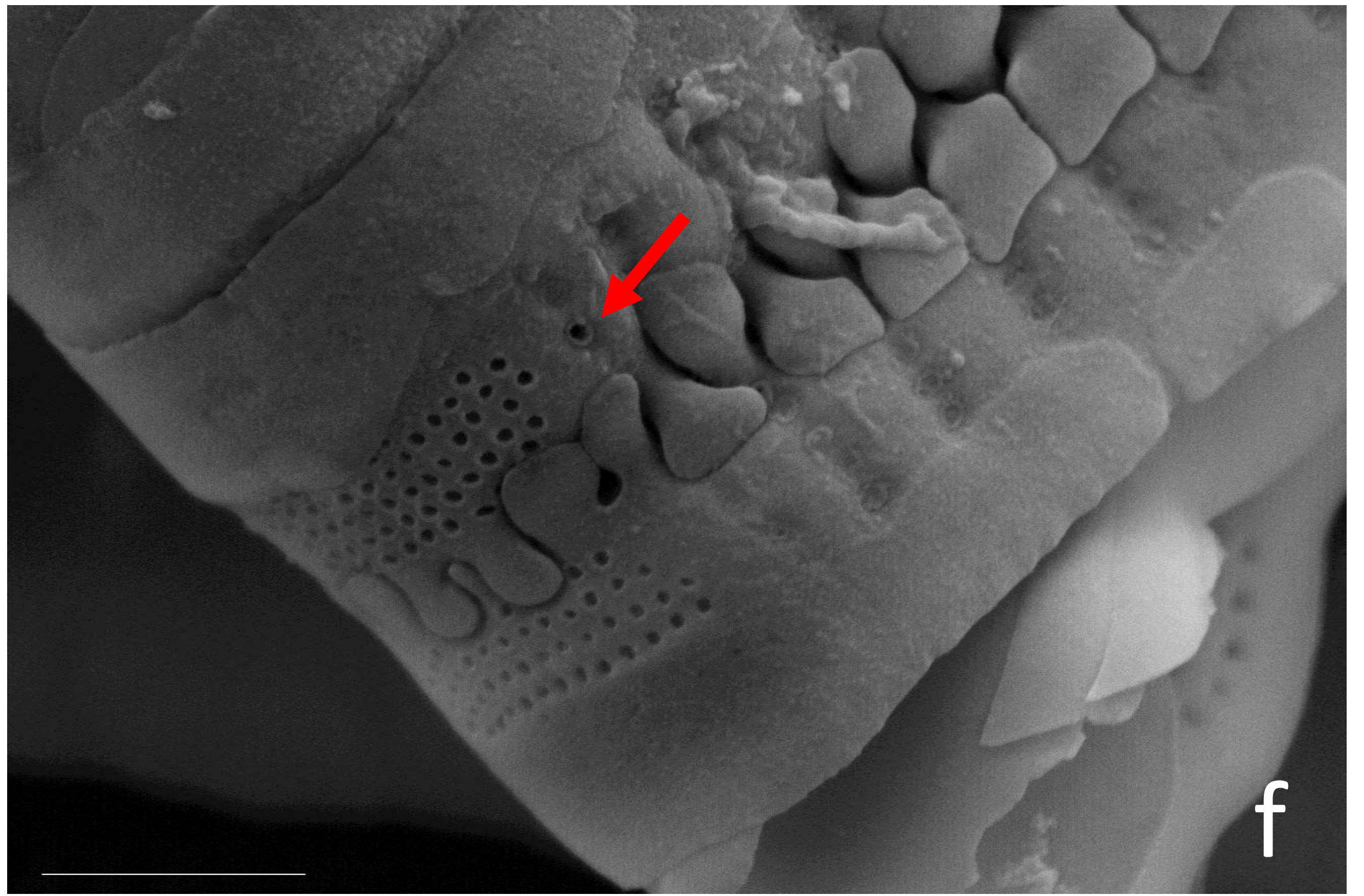
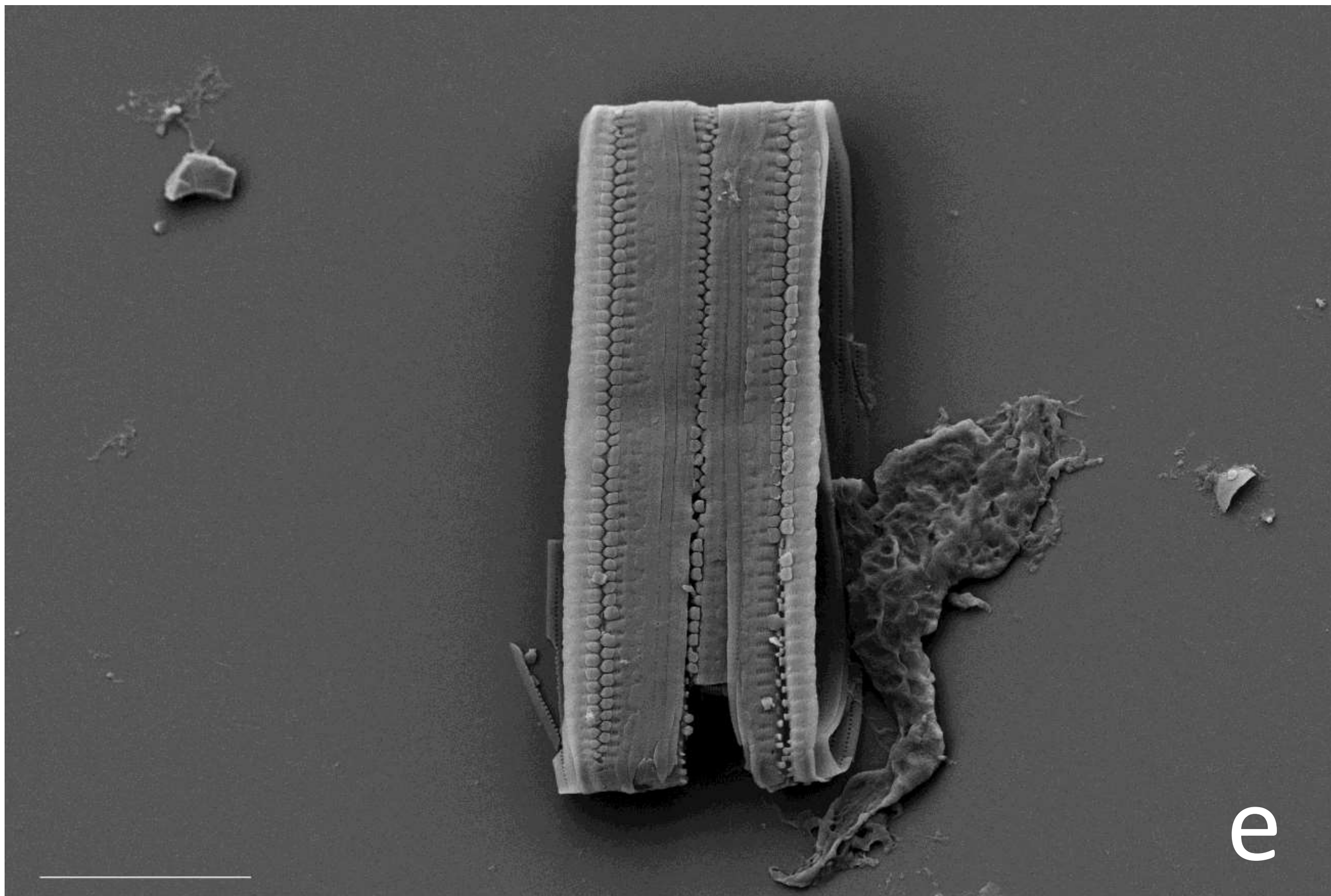
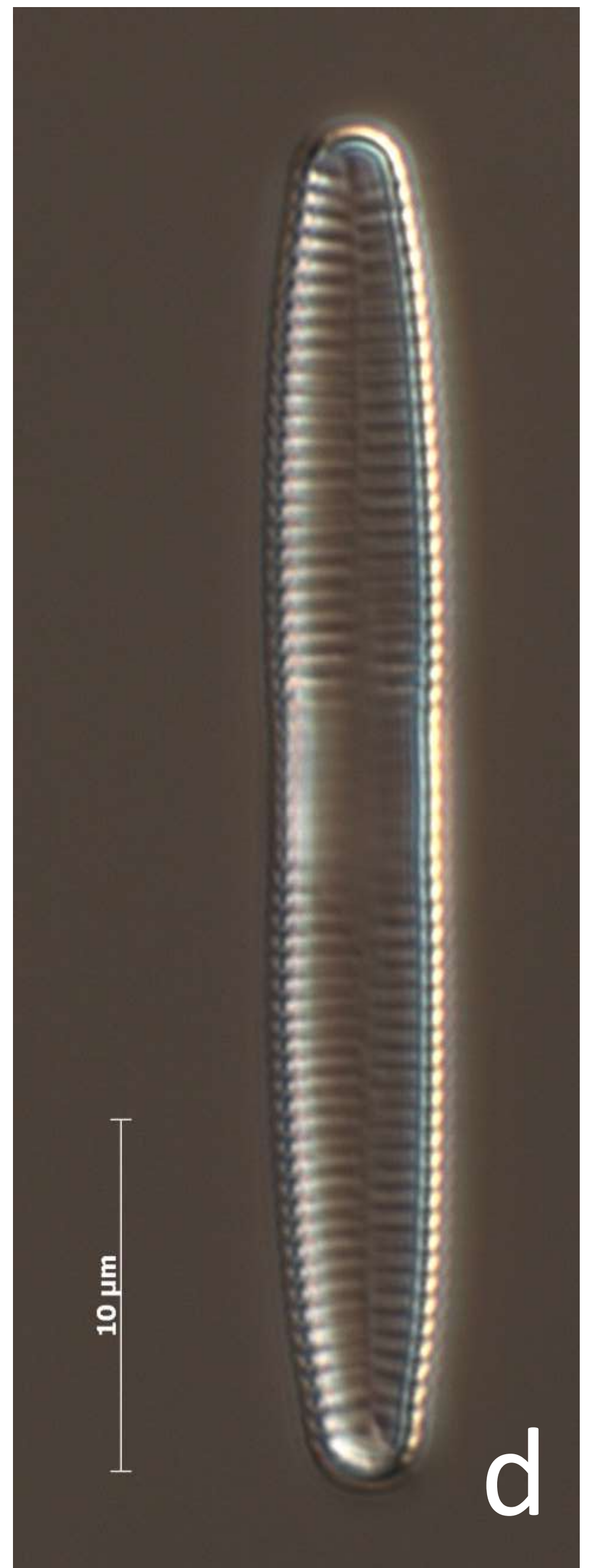
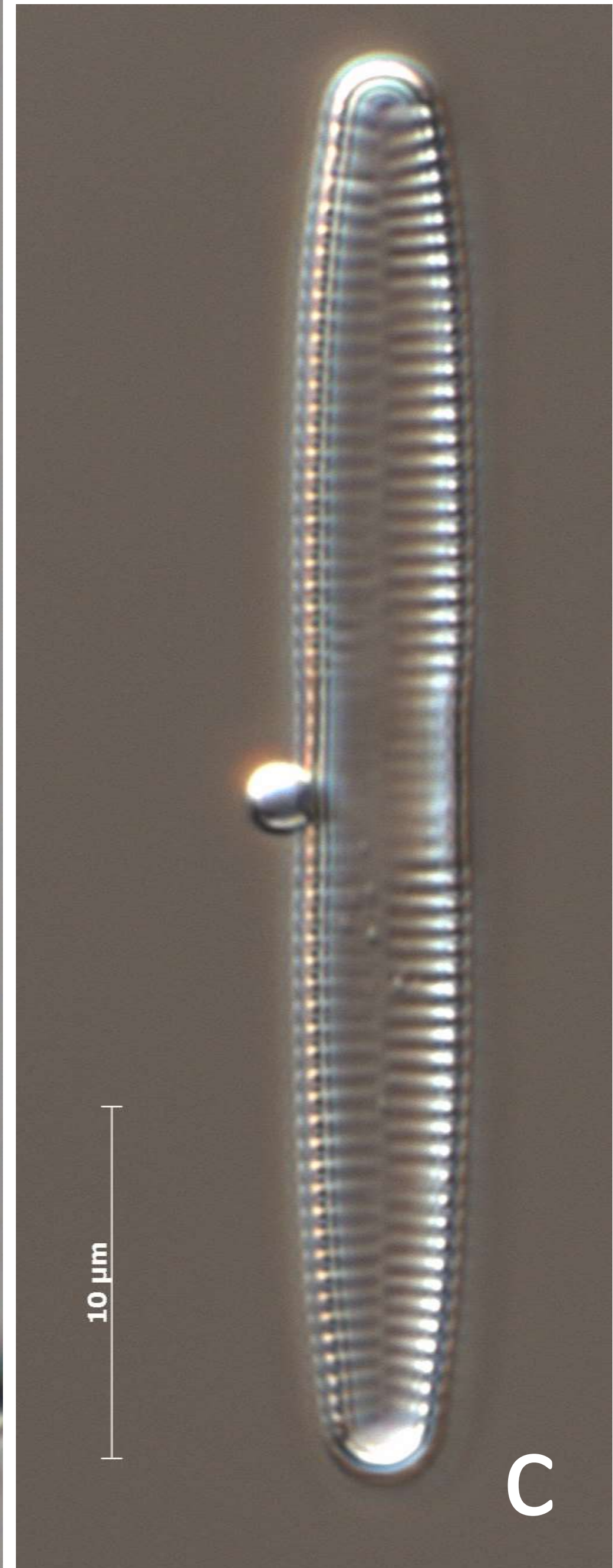
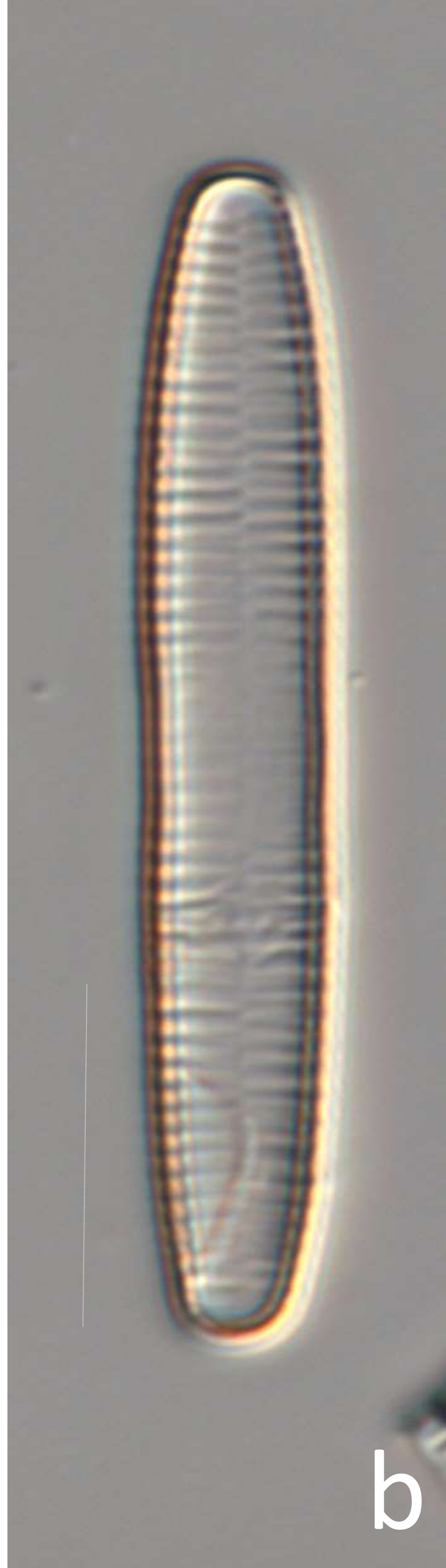
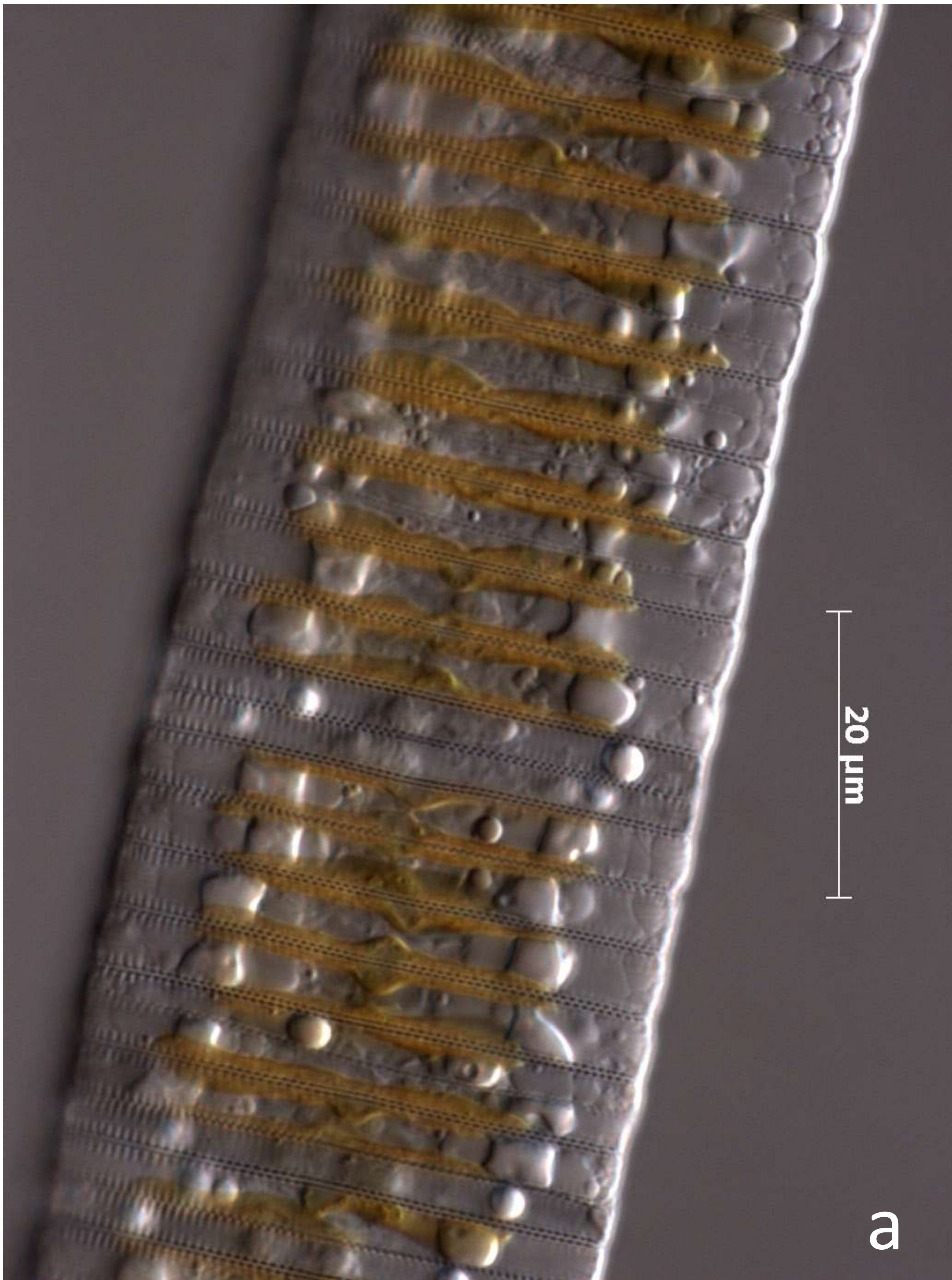


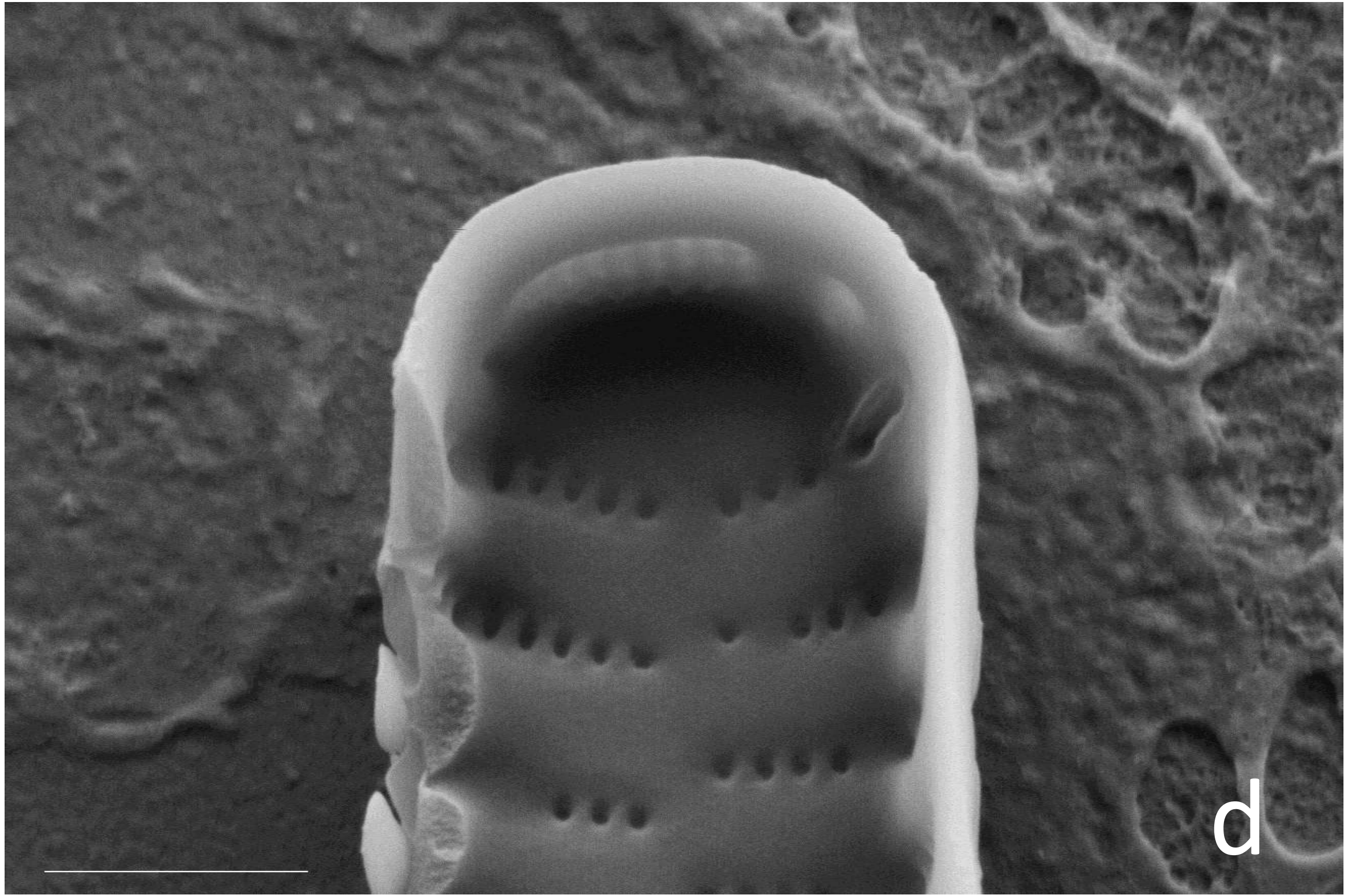
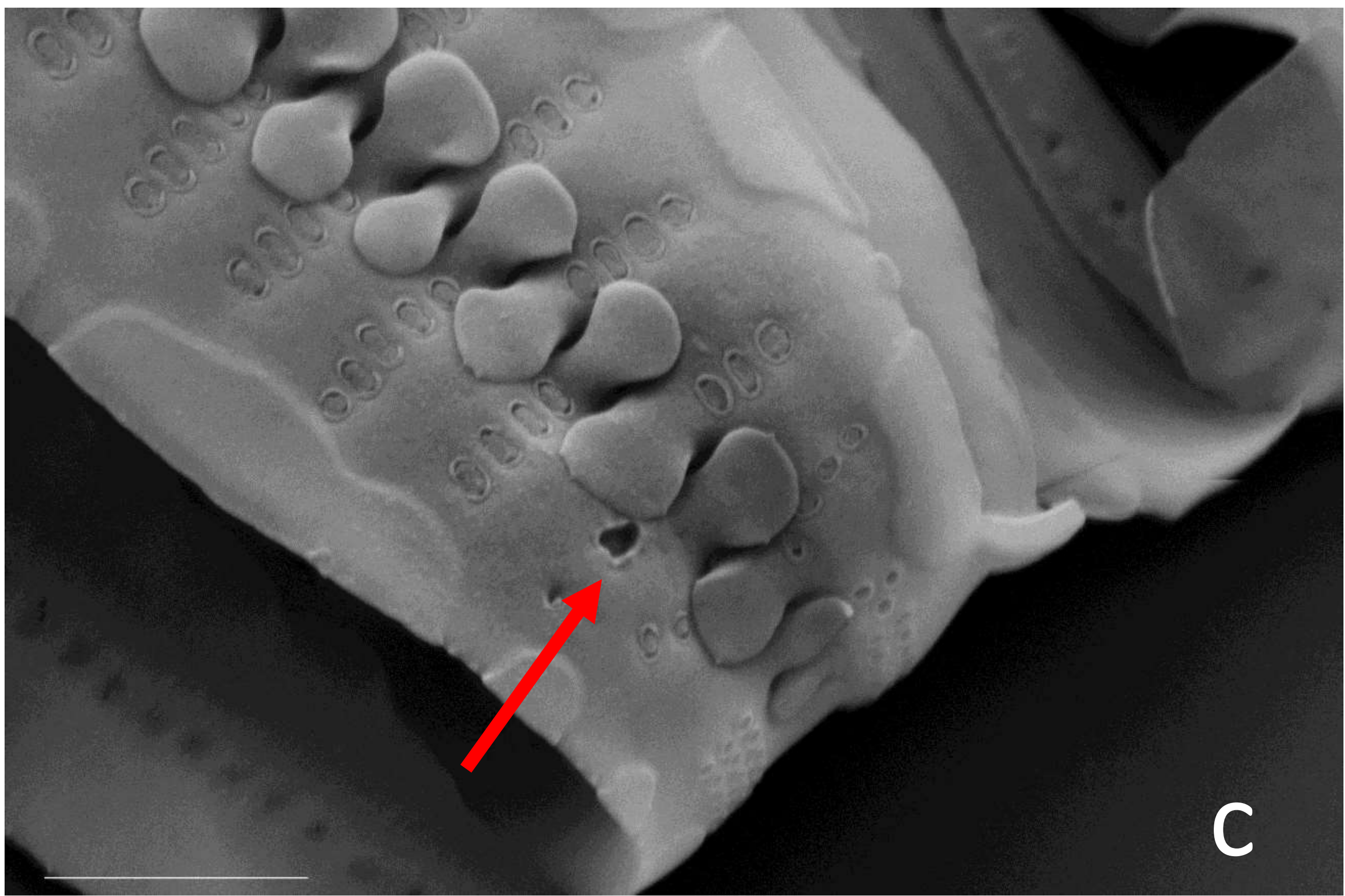
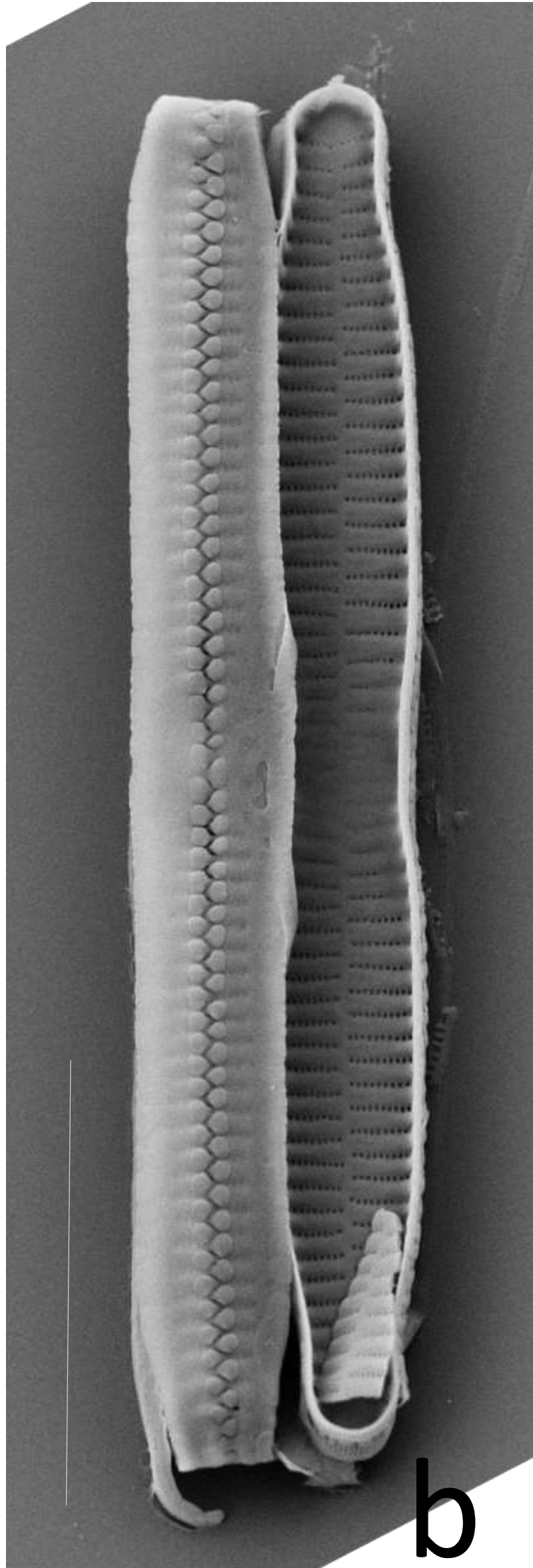
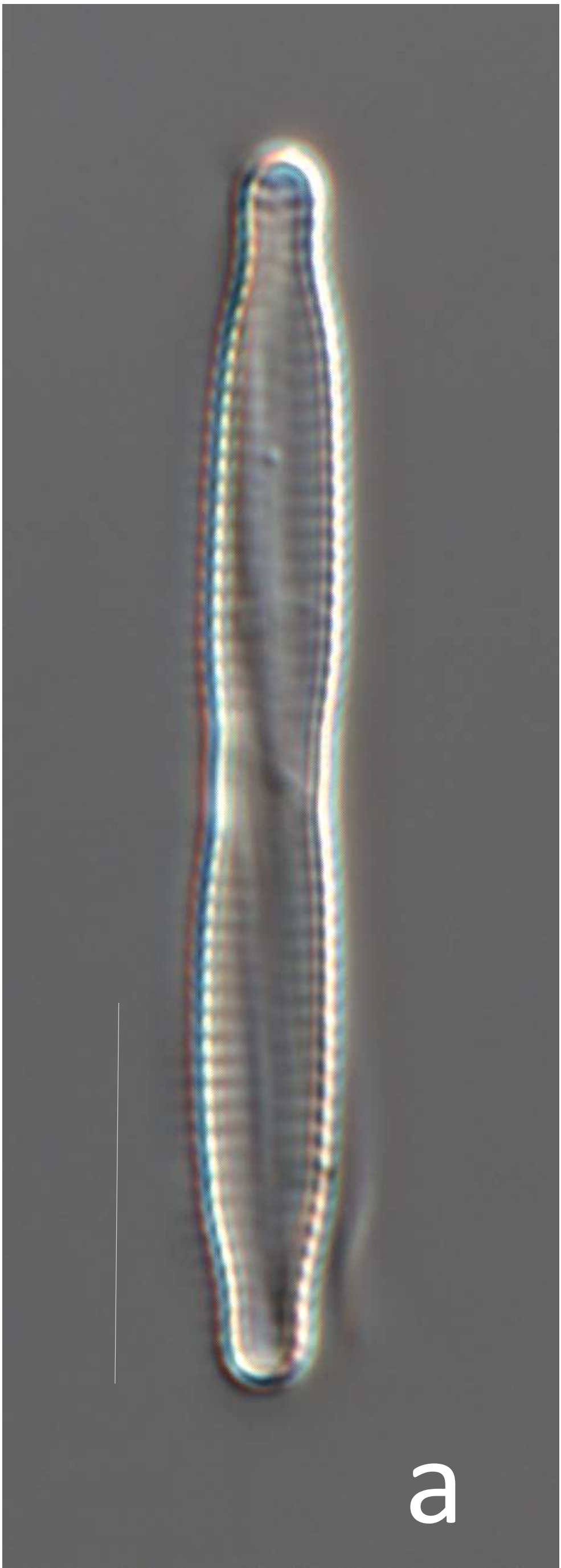


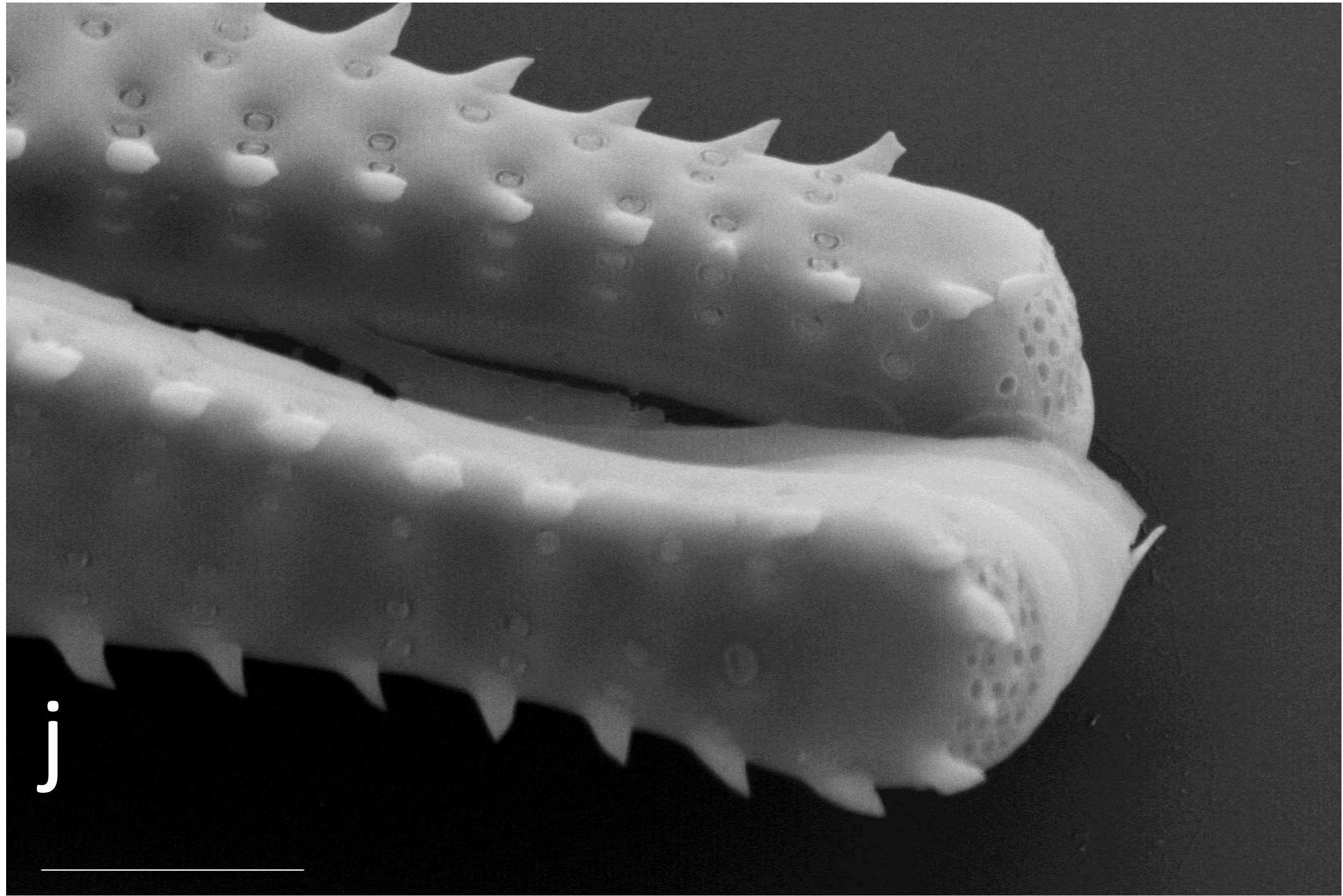
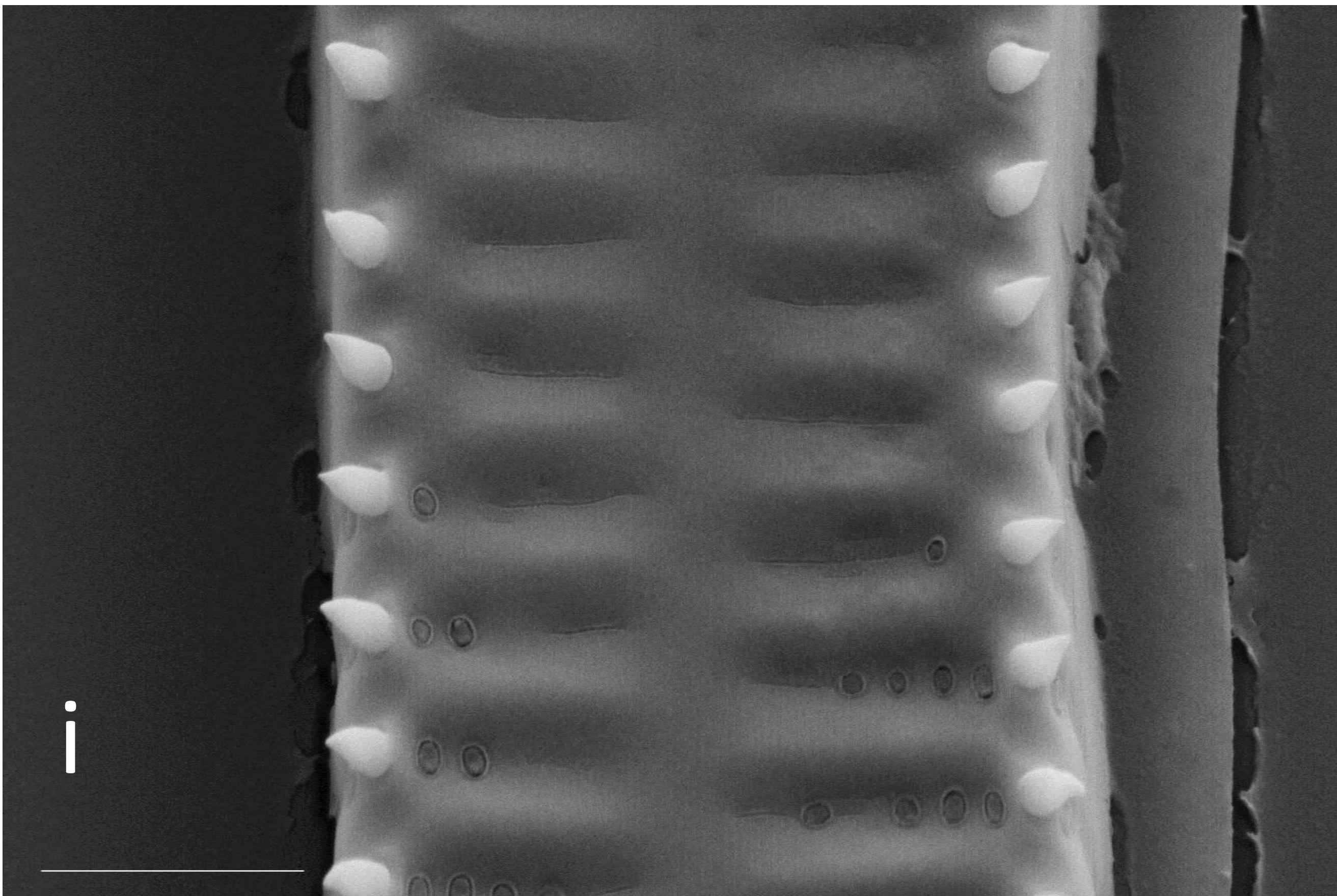
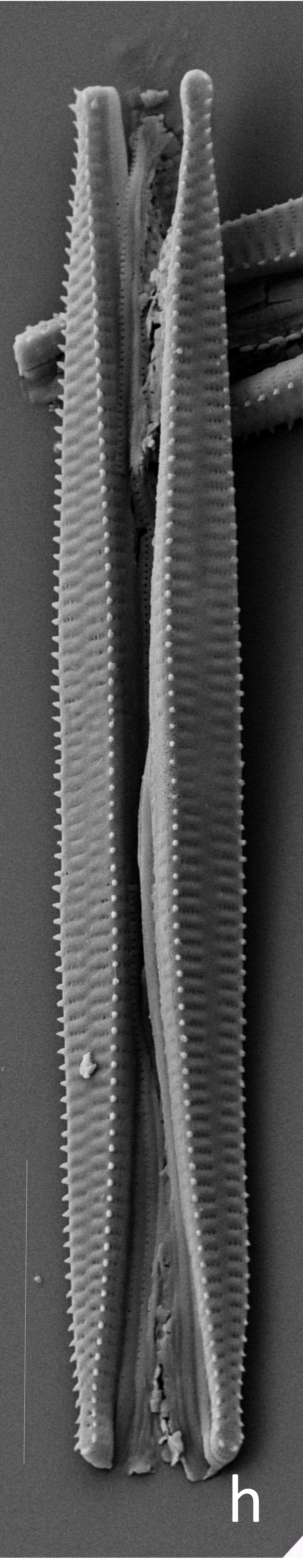
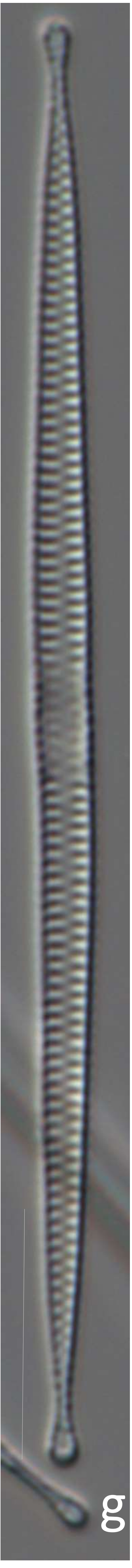
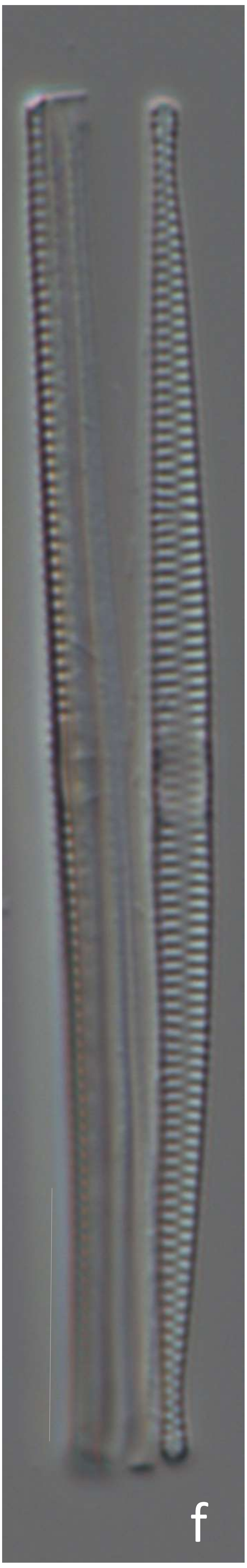
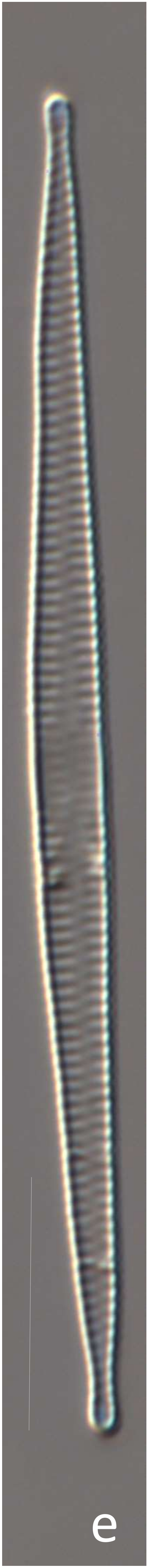
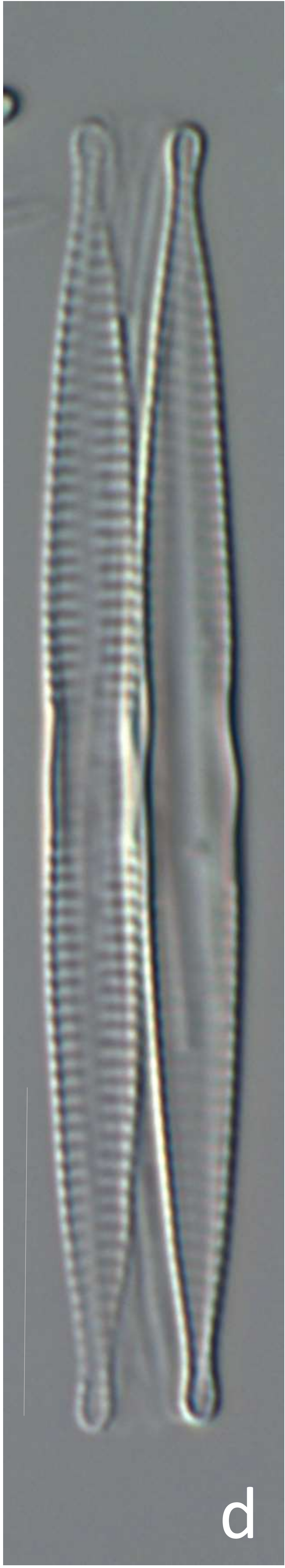
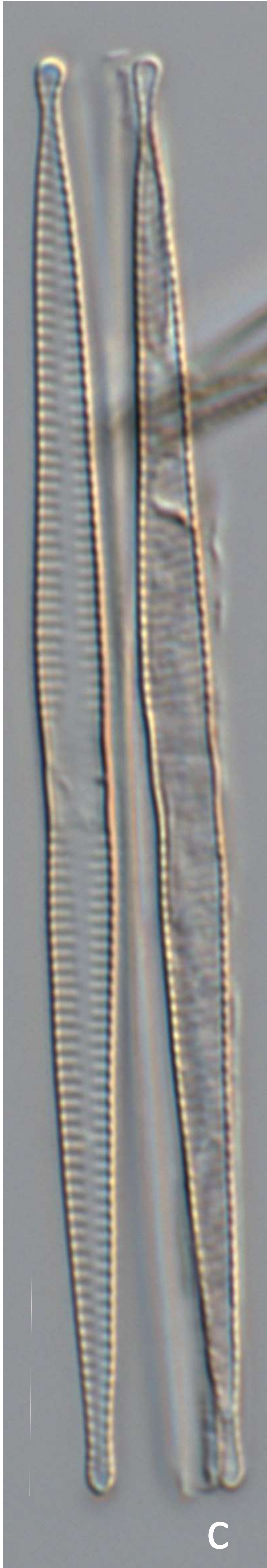
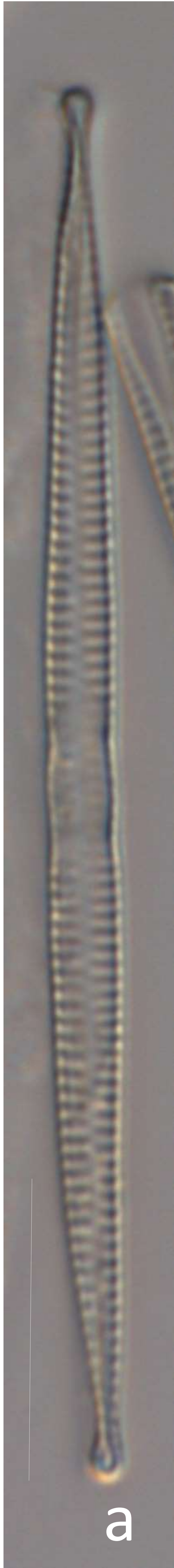


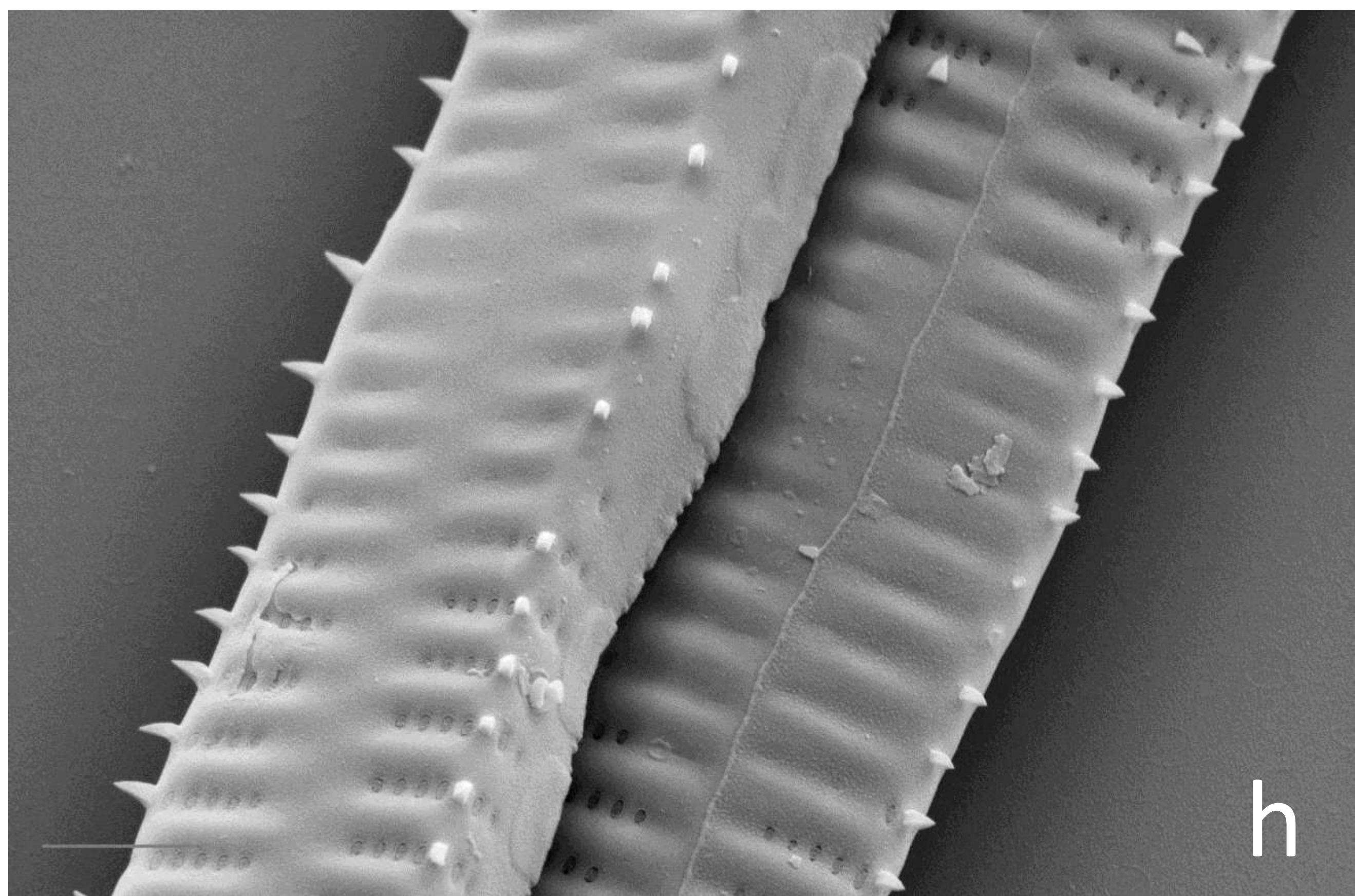
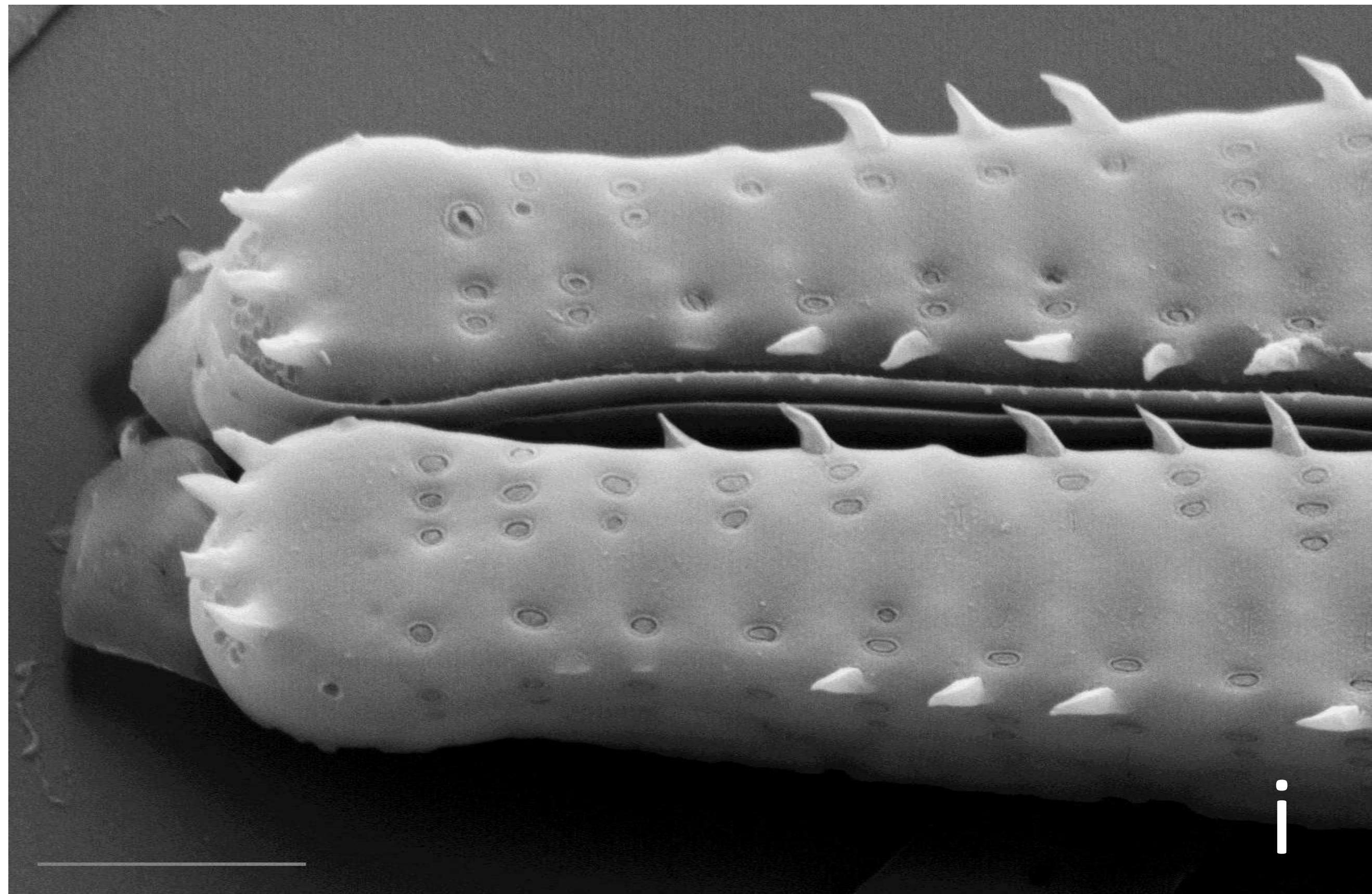
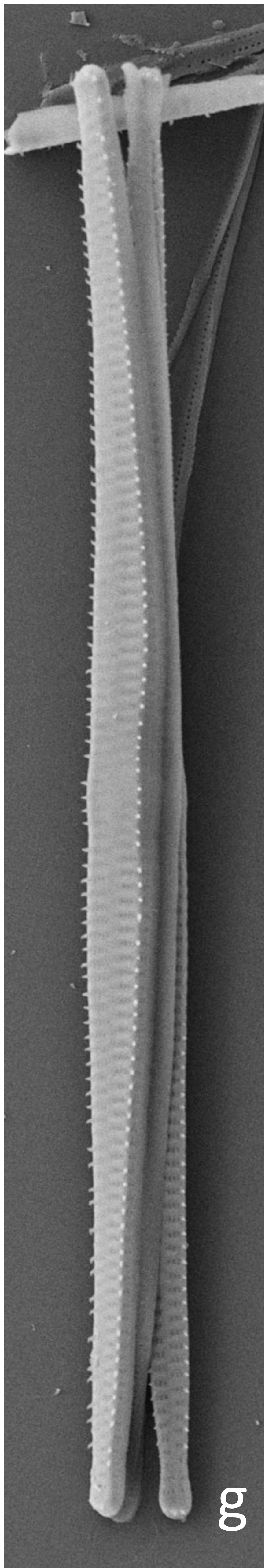
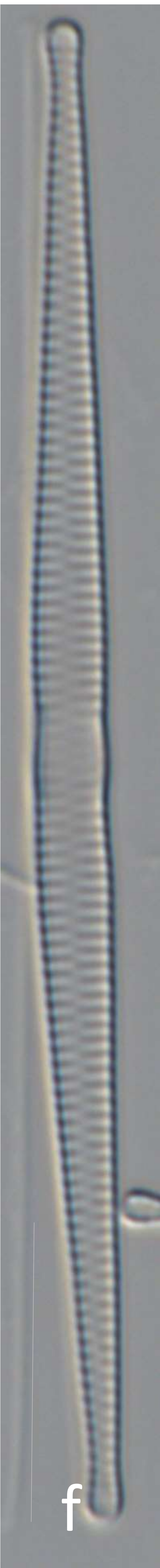
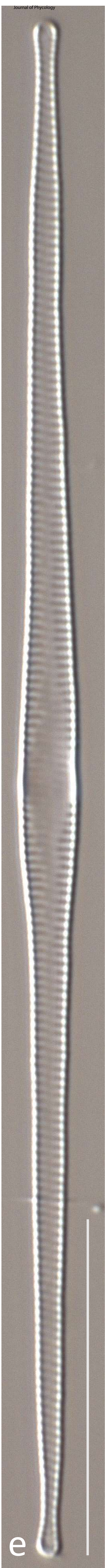
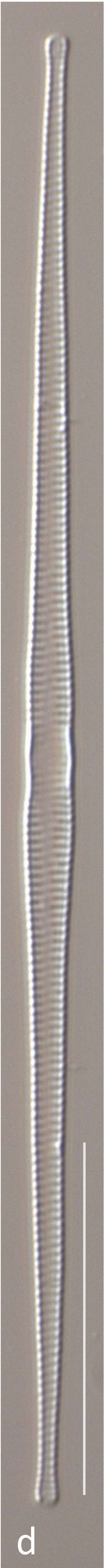
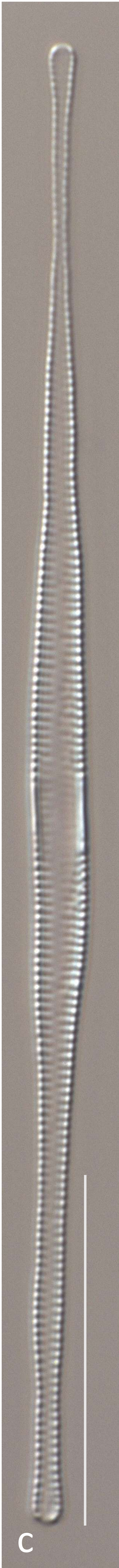
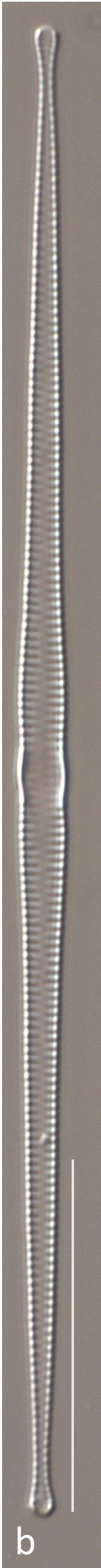












Clone ID	Voucher slide	original name	operational name
005FraP02	BC0005	<i>Fragilaria gracilis</i>	FGRA1
009FraP02	BC0009	<i>Fragilaria pararumpens</i>	FCRNAPA
012FraP02	BC0012	<i>Fragilaria gracilis</i>	FGRA1
015FraP02	BC0015	<i>Fragilaria</i> sp	FTNS
018FraP02	BC0018	<i>Fragilaria gracilis</i>	FGRA1
024SynP02	BC0024	<i>Ulnaria acus</i>	ULNA
028FraP02	BC0028	<i>Fragilaria gracilis</i>	FGRA1
032FraP02	BC0032	<i>Fragilaria</i> sp	FTEN2
033SynP02	BC0033	<i>Ulnaria acus</i>	ULNA
034FraP03	BC0034	<i>Fragilaria gracilis</i>	FGRA1
038FraP04	BC0038	<i>Fragilaria perminuta</i>	FPEM
041SynP04	BC0041	<i>Fragilaria vaucheriae</i>	FVAU
042SynP04	BC0042	<i>Fragilaria vaucheriae</i>	FCAP2
043SynP04	BC0043	<i>Fragilaria perminuta</i>	FPEM
046SynP04	BC0046	<i>Fragilaria vaucheriae</i>	FCAP2
047FraP04	BC0047	<i>Fragilaria gracilis</i>	FGRA1
048FraP04	BC0048	<i>Fragilaria pararumpens</i>	FCRNAPA
054SynP04	BC0054	<i>Fragilaria</i> cf. <i>pararumpens</i>	FCAP2
056FraP04	BC0056	<i>Fragilaria pararumpens</i>	FCRNAPA
061SynP05	BC0061	<i>Ulnaria ulna</i>	ULNA
085FraP07	BC0085	<i>Fragilaria gracilis</i>	FGRA1
091SynP07	BC0091	<i>Ulnaria ulna</i>	ULNA
121FraB01	BC0121	<i>Fragilaria gracilis</i>	FGRA2
135FraB03	BC0135	<i>Fragilaria gracilis</i>	FGRA1
139FraB04	BC0139	<i>Fragilaria gracilis</i>	FGRA2
142FraB04	BC0142	<i>Fragilaria gracilis</i>	FGRA2
143FraB04	BC0143	<i>Fragilaria gracilis</i>	FGRA1
148FraB04	BC0148	<i>Fragilaria gracilis</i>	FGRA2
151FraB04	BC0151	<i>Fragilaria gracilis</i>	FGRA2
153FraB04	BC0153	<i>Fragilaria gracilis</i>	FGRA2
154FraB04	BC0154	<i>Fragilaria</i> cf. <i>gracilis</i>	FGRA2
157FraB04	BC0157	<i>Fragilaria pararumpens</i>	FGRA2
158FraB04	BC0158	<i>Fragilaria gracilis</i>	FGRA2
162FraB05	BC0162	<i>Fragilaria gracilis</i>	FGRA2
166FraB05	BC0166	<i>Fragilaria gracilis</i>	FGRA2
167FraB05	BC0167	<i>Fragilaria gracilis</i>	FGRA2
169FraB05	BC0169	<i>Fragilaria gracilis</i>	FGRA2
170FraB05	BC0170	<i>Fragilaria gracilis</i>	FGRA2
171FraB05	BC0171	<i>Fragilaria vaucheriae</i>	FCAP2
173FraB05	BC0173	<i>Fragilaria gracilis</i>	FGRA2
174FraB05	BC0174	<i>Fragilaria gracilis</i>	FGRA2
175FraB05	BC0175	<i>Fragilaria gracilis</i>	FGRA2
177FraB05	BC0177	<i>Fragilaria gracilis</i>	FGRA2
178FraB05	BC0178	<i>Fragilaria gracilis</i>	FGRA2
185FraB06	BC0185	<i>Fragilaria gracilis</i>	FGRA1
191FraB06	BC0191	<i>Fragilaria pararumpens</i>	FCRNAPA
194FraB06	BC0194	<i>Fragilaria crotonensis</i>	FCRNAPA
195FraB07	BC0195	<i>Fragilaria pararumpens</i>	FCRNAPA
196FraB07	BC0196	<i>Fragilaria pararumpens</i>	FCRNAPA

199FraB07	BC0199	Fragilaria pararumpens	FCRNAPA
203FraB07	BC0203	Fragilaria pararumpens	FCRNAPA
205FraB07	BC0205	Fragilaria pararumpens	FCRNAPA
215UlnB08	BC0215	Ulnaria acus	ULNA
220UlnB08	BC0220	Ulnaria ulna	ULNA
241FraB08	BC0241	Fragilaria gracilis	FGRA2
243FraB09	BC0243	Fragilaria pararumpens	FCRNAPA
245FraB09	BC0245	Fragilaria pararumpens	FCRNAPA
246FraB09	BC0246	Fragilaria pararumpens	FCRNAPA
247FraB09	BC0247	Fragilaria pararumpens	FCRNAPA
248FraB09	BC0248	Fragilaria pararumpens	FCRNAPA
249FraB09	BC0249	Fragilaria pararumpens	FCRNAPA
266FraB10	BC0266	Fragilaria gracilis	FGRA2
319FraW02	BC0319	Fragilaria gracilis	FGRA1
343FraT01	BC0343	Fragilaria sp	FTEN
344UlnT01	BC0344	Ulnaria ulna	ULNA
358UlnW01	BC0358	Ulnaria ulna	ULNA
378FraB10	BC0378	Fragilaria gracilis	FGRA2
379FraB10	BC0379	Fragilaria gracilis	FGRA2
380FraB10	BC0380	Fragilaria gracilis	FGRA2
381FraB10	BC0381	Fragilaria gracilis	FGRA2
382FraB10	BC0382	Fragilaria gracilis	FGRA2
383FraB10	BC0383	Fragilaria gracilis	FGRA2
385FraB10	BC0385	Fragilaria gracilis	FGRA2
387FraB10	BC0387	Fragilaria gracilis	FGRA2
388FraB10	BC0388	Fragilaria gracilis	FGRA2
404FraB10	BC0404	Fragilaria gracilis	FGRA2
409FraB10	BC0409	Fragilaria gracilis	FGRA2
410FraB10	BC0410	Fragilaria gracilis	FGRA2
412FraB10	BC0412	Fragilaria gracilis	FGRA2
414FraB10	BC0414	Fragilaria gracilis	FGRA2
435FraT01	BC0435	Fragilaria vaucheriae	FCAP2
467FraR03	BC0467	Fragilaria gracilis	FGRA1
488FraR03	BC0488	Fragilaria gracilis	FGRA1
511FraK01	BC0511	Fragilaria gracilis	FGRA2
513FraK01	BC0513	Fragilaria capucina	FCAP1
514FraK01	BC0514	Fragilaria capucina	FCAP1
515FraK01	BC0515	Fragilaria gracilis	FGRA2
516FraK01	BC0516	Fragilaria gracilis	FGRA2
517FraK01	BC0517	Fragilaria gracilis	FGRA2
551FraK01	BC0551	Fragilaria gracilis	FGRA1
555FraK01	BC0555	Fragilaria gracilis	FGRA2
575FraK01	BC0575	Fragilaria rumpens	FTEN
591FraK01	BC0591	Fragilaria gracilis	FGRA2
613FraP11	BC0613	Fragilaria sp	FTEN
621FraP11	BC0621	Fragilaria capucina	FCAP1
622FraP11	BC0622	Fragilaria gracilis	FGRA1
625FraP11	BC0625	Fragilaria rumpens	FTEN
643FraK06	BC0643	Fragilaria gracilis	FGRA1
653FraK08	BC0653	Fragilaria mesolepta	FCRNAPA

657UlnK08	BC0657	Ulnaria ulna	ULNA
662FraK09	BC0662	Fragilaria sp	FTEN
665SynK09	BC0665	Ulnaria ulna	ULNA
668SynK09	BC0668	Ulnaria acus	ULNA
698UlnK10	BC0698	Ulnaria ulna	ULNA
702UlnK11	BC0702	Ulnaria ulna	ULNA
706SynK11	BC0706	Ulnaria acus	ULNA
720UlnK13	BC0720	Ulnaria ulna	ULNA
726FraB12	BC0726	Fragilaria sp	FTEN
732FraB12	BC0732	Fragilaria tenera	FTEN
734FraB13	BC0734	Fragilaria gracilis	FGRA2
741FraB13	BC0741	Fragilaria gracilis	FGRA2
743FraB14	BC0743	Fragilaria gracilis	FGRA2
746FraB14	BC0746	Fragilaria gracilis	FGRA2
791FraN01	BC0791	Fragilaria sp	FRAS
819FraN02	BC0819	Fragilaria pararumpens	FCRNAPA
820SynN05	BC0820	Ulnaria acus	ULNA
823SynN05	BC0823	Fragilaria pararumpens	FCRNAPA
836UlnN05	BC0836	Ulnaria ulna	ULNA
842FraN04	BC0842	Fragilaria gracilis	FGRA2
844FraN04	BC0844	Fragilaria gracilis	FGRA2
845FraN04	BC0845	Fragilaria gracilis	FGRA2
852FraN04	BC0852	Fragilaria gracilis	FGRA2
853FraN04	BC0853	Fragilaria pararumpens	FCRNAPA
854FraN04	BC0854	Fragilaria gracilis	FGRA2
855FraN04	BC0855	Fragilaria gracilis	FGRA1
856FraN04	BC0856	Fragilaria gracilis	FGRA2
881UlnR05	BC0881	Ulnaria ulna	ULNA
907FraR05	BC0907	Fragilaria gracilis	FGRA1
s0327	s0327	Fragilaria bidens	FRAS
AT-185Gel3	n/a	Fragilaria crotonensis	FCRNAPA
AT_185Gel3	n/a	Fragilaria crotonensis	FCRNAPA
AT-124.05b	n/a	Fragilaria vaucheriae	FRAS
CCAP1011/1	n/a	Centronella reicheltii	FCRNAPA
UTEXFD404	n/a	Ulnaria ulna	ULNA
TCC134	TCC134	Ulnaria acus	ULNA
TCC301	TCC301	Fragilaria crotonensis	FCRNAPA
TCC302	TCC302	Fragilaria crotonensis	FCRNAPA
TCC304	TCC304	Fragilaria crotonensis	FCRNAPA
TCC306	TCC306	Ulnaria ulna	ULNA
TCC365	TCC365	Fragilaria crotonensis	FCRNAPA
TCC367	TCC367	Fragilaria perminuta	FPEM
TCC520	TCC520	Ulnaria ulna	ULNA
TCC522	TCC522	Ulnaria ulna	ULNA
TCC541	TCC541	Fragilaria vaucheriae	FVAU
TCC547	TCC547	Fragilaria vaucheriae	FVAU
TCC553	TCC553	Fragilaria vaucheriae	FVAU
TCC558	TCC558	Fragilaria vaucheriae	FVAU
TCC559	TCC559	Fragilaria capucina	FRAS
TCC562	TCC562	Fragilaria capucina	FCRNAPA

TCC584	TCC584	<i>Ulnaria acus</i>	ULNA
TCC589	TCC589	<i>Fragilaria capucina</i>	FCRNAPA
TCC626	TCC626	<i>Ulnaria ulna</i>	ULNA
TCC633	TCC633	<i>Ulnaria ulna</i>	ULNA
TCC634	TCC634	<i>Ulnaria ulna</i>	ULNA
TCC635	TCC635	<i>Ulnaria ulna</i>	ULNA
TCC654	TCC654	<i>Ulnaria ulna</i>	ULNA
TCC656	TCC656	<i>Ulnaria ulna</i>	ULNA
TCC662	TCC662	<i>Fragilaria rumpens</i>	FVAU
TCC666	TCC666	<i>Fragilaria rumpens</i>	FGRA1
TCC669	TCC669	<i>Fragilaria rumpens</i>	FGRA2
TCC670	TCC670	<i>Ulnaria ulna</i>	ULNA
TCC671	TCC671	<i>Fragilaria rumpens</i>	FGRA2
TCC673	TCC673	<i>Fragilaria rumpens</i>	FGRA2
TCC677	TCC677	<i>Fragilaria rumpens</i>	FGRA1
TCC681	TCC681	<i>Fragilaria rumpens</i>	FVAU
TCC682	TCC682	<i>Fragilaria rumpens</i>	FCAP1
TCC686	TCC686	<i>Fragilaria rumpens</i>	FGRA1
TCC695	TCC695	<i>Ulnaria ulna</i>	ULNA
TCC699	TCC699	<i>Ulnaria ulna</i>	ULNA
TCC705	TCC705	<i>Fragilaria capucina</i>	FRAS
TCC716	TCC716	<i>Ulnaria ulna</i>	ULNA
TCC722	TCC722	<i>Fragilaria rumpens</i>	FGRA1
TCC728	TCC728	<i>Fragilaria rumpens</i>	FGRA1
TCC729	TCC729	<i>Fragilaria rumpens</i>	FGRA1
TCC743	TCC743	<i>Fragilaria perminuta</i>	FPEM
TCC747	TCC747	<i>Fragilaria perminuta</i>	FPEM
TCC752	TCC752	<i>Fragilaria perminuta</i>	FPEM
TCC7a	TCC7a	<i>Fragilaria vaucheriae</i>	FCRNAPA
TCC829	TCC829	<i>Fragilaria perminuta</i>	FPEM
TCC862	TCC862	<i>Fragilaria cf. nanoides</i>	FCRNAPA
TCC863	TCC863	<i>Fragilaria cf. nanoides</i>	FCRNAPA
TCC865	TCC865	<i>Fragilaria perminuta</i>	FPEM
TCC866	TCC866	<i>Fragilaria perminuta</i>	FPEM
TCC867	TCC867	<i>Fragilaria tenuistriata</i>	FTNS
TCC868	TCC868	<i>Fragilaria tenuistriata</i>	FTNS
TCC869	TCC869	<i>Fragilaria gracilis</i>	FGRA1
TCC870	TCC870	<i>Fragilaria cf. nanoides</i>	FTEN2
TCC871	TCC871	<i>Fragilaria cf. nanoides</i>	FTEN2
TCC873	TCC873	<i>Fragilaria perminuta</i>	FPEM
TCC874	TCC874	<i>Fragilaria perminuta</i>	FPEM
TCC877	TCC877	<i>Fragilaria capucina</i> var. <i>capucina</i>	FCAP2
TCC878	TCC878	<i>Fragilaria cf. nanoides</i>	FCRNAPA
TCC882	TCC882	<i>Fragilaria perminuta</i>	FPEM
TCC887	TCC887	<i>Fragilaria capucina</i> var. <i>capucina</i>	FCAP2

Fragilaria nanoides

Fragilaria nanoides
Fragilaria pararumpens
Fragilaria nanoides
Ulnaria ulna
Ulnaria ulna
Fragilaria gracilis
Fragilaria nanoides
Fragilaria nanoides
Fragilaria nanoides
Fragilaria cf. nanoides
Fragilaria cf. nanoides
Fragilaria cf. nanoides
Fragilaria gracilis
Fragilaria gracilis
Fragilaria tenera
Ulnaria ulna
Ulnaria ulna
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
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Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria joachimii
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria heatherae
Fragilaria heatherae
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria tenera
Fragilaria gracilis
Fragilaria tenera
Fragilaria heatherae
Fragilaria gracilis
Fragilaria tenera
Fragilaria gracilis
Fragilaria mesolepta

Ulnaria ulna
Fragilaria tenera
Ulnaria ulna
Ulnaria acus
Ulnaria ulna
Ulnaria ulna
Ulnaria acus
Ulnaria ulna
Fragilaria tenera
Fragilaria tenera
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria sp.
Fragilaria cf. pararumpens
Ulnaria acus
Fragilaria pararumpens
Ulnaria ulna
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria pararumpens
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Ulnaria ulna
Fragilaria gracilis
Fragilaria bidens
Fragilaria crotonensis
Fragilaria crotonensis
Fragilaria cf. vaucheriae
Centronella reicheltii
Ulnaria ulna
Ulnaria acus
Fragilaria crotonensis
Fragilaria crotonensis
Fragilaria crotonensis
Ulnaria ulna
Fragilaria crotonensis
Fragilaria perminuta
Ulnaria ulna
Ulnaria ulna
Fragilaria agnesiae
Fragilaria agnesiae
Fragilaria agnesiae
Fragilaria agnesiae
Fragilaria cf. capucina
Fragilaria cf. capucina

Ulnaria acus
Fragilaria cf. capucina
Ulnaria ulna
Ulnaria ulna
Ulnaria ulna
Ulnaria ulna
Ulnaria ulna
Ulnaria ulna
Fragilaria agnesiae
Fragilaria cf. gracilis
Fragilaria cf. rumpens
Ulnaria ulna
Fragilaria gracilis
Fragilaria gracilis
Fragilaria cf. gracilis
Fragilaria agnesiae
Fragilaria heatherae
Fragilaria cf. gracilis
Ulnaria ulna
Ulnaria ulna
Fragilaria sp.
Ulnaria ulna
Fragilaria gracilis
Fragilaria gracilis
Fragilaria gracilis
Fragilaria perminuta
Fragilaria perminuta
Fragilaria perminuta
Fragilaria sp.
Fragilaria perminuta
Fragilaria cf. nanoides
Fragilaria cf. nanoides
Fragilaria perminuta
Fragilaria perminuta
Fragilaria subconstricta
Fragilaria subconstricta
Fragilaria gracilis
Fragilaria sp. 1
Fragilaria sp. 1
Fragilaria perminuta
Fragilaria perminuta
Fragilaria joachimii
Fragilaria cf. nanoides
Fragilaria perminuta
Fragilaria joachimii

Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Water of Leith, Currie, Edinburgh	United Kingdom
River Tay, near Aberfeldy, Perth & Kinross	United Kingdom
River Tay, near Aberfeldy, Perth & Kinross	United Kingdom
Water of Leith, Currie, Edinburgh	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
Allt a 'Bhalachain, Argyll & Bute	United Kingdom
River Tay, near Aberfeldy, Perth & Kinross	United Kingdom
Royal Botanic Garden Edinburgh, pond	United Kingdom
Royal Botanic Garden Edinburgh, pond	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
Euden Beck	United Kingdom
River Tay, Pitlochry, Perth & Kinross	United Kingdom
River Tay, Pitlochry, Perth & Kinross	United Kingdom
River Tay, Pitlochry, Perth & Kinross	United Kingdom
River Tay, Pitlochry, Perth & Kinross	United Kingdom
Cheriton Stream, Cheriton	United Kingdom
Pillhill Brook, Upper Clatford (112278)	United Kingdom

Pillhill Brook, Upper Clatford (112278)	United Kingdom
River Anton, Andover, "KFC"	United Kingdom
River Anton, Andover, "KFC"	United Kingdom
River Anton, Andover, "KFC"	United Kingdom
Lambourn, Bagnor (112280)	United Kingdom
River Kennet, Stitchcombe Mill	United Kingdom
River Kennet, Stitchcombe Mill	United Kingdom
River Wylfe, Henford Marsh	United Kingdom
Inveruglas Water, by Ben Vane, Argyll & Bute	United Kingdom
Inveruglas Water, by Ben Vane, Argyll & Bute	United Kingdom
Inveruglas Water, by Ben Vane, Argyll & Bute	United Kingdom
Inveruglas Water, by Ben Vane, Argyll & Bute	United Kingdom
Allt Coiregrogain, by Ben Vane, Argyll & Bute	United Kingdom
Allt Coiregrogain, by Ben Vane, Argyll & Bute	United Kingdom
Wooler Water near Wooler, Northumbria	United Kingdom
Wooler Water near Wooler, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Harthope Burn, Northumbria	United Kingdom
Royal Botanic Garden Edinburgh, pond	United Kingdom
Royal Botanic Garden Edinburgh, pond	United Kingdom
Okinoshima Island, Shimane Prefecture	Japan
Wümme River, Schleswig-Holstein	Germany
Wümme River, Schleswig-Holstein	Germany
Zarrentiner Becken Lake, Mecklenburg-Vorpommern	Germany
Plußsee, Rathjensdorf, Plön	Germany
Minnesota	USA
Lac d'Annecy	France
Lac du Bourget	France
Lac du Bourget	France
Lac du Bourget	France
Lac du Bourget	France
Lac Léman - SHL2	France
Lac Léman, Port de l'INRA	France
Ile de La Réunion rivière de des Galets site Marla	France
Ile de La Réunion rivière de des Galets site Marla	France
Ile de La Réunion rivière de Bras Caverne	France
Ile de La Réunion rivière de Langevin, grand Galet	France
Ile de La Réunion rivière de Langevin, grand Galet	France
Ile de La Réunion rivière de Langevin site amont prise EDF	France
Ile de La Réunion rivière de Sainte Suzanne	France
Ile de La Réunion rivière de Sainte Suzanne	France

rivière la Moselle à Bainville aux Miroirs	France
Canal de Nantes à Brest à Nort-sur-Erdre	France
rivière Le Gier à Givors	France
rivière Eischbaach à Boevange/attert	Luxembourg
rivière Eischbaach à Boevange/attert	Luxembourg
rivière Eischbaach à Boevange/attert	Luxembourg
rivière Alzette à Walfer-Steinsel	Luxembourg
rivière Attert à Colmar-Berg	Luxembourg
rivière Our à Vianden	Luxembourg
rivière Sûre à Camping Heiderscheidergrund	Luxembourg
Trentino Canal à Vérone	Italy
Trentino rivière de Avisio à Lavis	Italy
Trentino rivière de Brusago	Italy
Trentino rivière de Brusago	Italy
Trentino rivière de Brusago	Italy
Trentino rivière de Regnana à Amont de Bedollo	Italy
Trentino rivière de Regnana à Amont de Bedollo	Italy
Piemonte rivière de Rocciamelone à Foresto (Bussoleno)	Italy
Trento Torrente à Fersina	Italy
rivière de Carrión à Embalse de Compuerto	Spain
rivière de Valdavia à Osorno	Spain
rivière de Yuso à Boca de Huérgano	Spain
Rib. De Seixe à Zambujeira De Baixo	Portugal
Rib. De Seixe à Foz Do Arroio	Portugal
Rib. De Seixe à Foz Do Arroio	Portugal
Lac Léman, estuaire du Foron	France
Lac Léman, estuaire du Foron	France
Lac Léman, parc de Rovorée	France
rivière La Loue (Jura)	France
Lac Léman, Port de l'INRA, France	France
Siggeforasjön lake near Norrtälje city	Sweden
Siggeforasjön lake near Norrtälje city	Sweden
Erken lake near Norrtälje city	Sweden
Erken lake near Norrtälje city	Sweden
Järsöströmmen river near Norrtälje city	Sweden
Järsöströmmen river near Norrtälje city	Sweden
Siggeforasjön lake near Norrtälje city	Sweden
Siggeforasjön lake near Norrtälje city	Sweden
Siggeforasjön lake near Norrtälje city	Sweden
Erken lake near Norrtälje city	Sweden
Erken lake near Norrtälje city	Sweden
Norrtäljeån river near Norrtälje city	Sweden
Siggeforasjön lake near Norrtälje city	Sweden
Erken lake near Norrtälje city	Sweden
Broströmmen river near Norrtälje city	Sweden

Sampling site	Sampling site	Culture autho	Sampling date	Sequence ID	rbcl sequenc
55.844216	-3.311093	D. Mann, M. K	2012/05/19	005FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	009FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	012FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	015FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	018FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	024SynP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	028FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	032FraP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	033SynP02	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	034FraP03	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	038FraP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	041SynP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	042SynP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	043SynP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	046SynP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	047FraP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	048FraP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	054SynP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	056FraP04	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	061SynP05	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	085FraP07	ATGTCTCAA1
55.844216	-3.311093	D. Mann, M. K	2012/05/19	091SynP07	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	121FraB01	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	135FraB03	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	139FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	142FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	143FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	148FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	151FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	153FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	154FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	157FraB04	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	158FraB04	ATGTCTCAA1
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56.206051	-4.790649	D. Mann, M. K	2012/05/19	169FraB05	ATGTCTCAA1
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56.206051	-4.790649	D. Mann, M. K	2012/05/19	173FraB05	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	174FraB05	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	175FraB05	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	177FraB05	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	178FraB05	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	185FraB06	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	191FraB06	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/26	194FraB06	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	195FraB07	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19	196FraB07	ATGTCTCAA1

56.206051	-4.790649	D. Mann, M. K	2012/05/19 199FraB07	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 203FraB07	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 205FraB07	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 215UlnB08	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 220UlnB08	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 241FraB08	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 243FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 245FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 246FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 247FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 248FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 249FraB09	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 266FraB10	ATGTCTCAA1
55.895467	-3.308024	D. Mann, M. K	2012/05/19 319FraW02	ATGTCTCAA1
56.6177	-3.8837	D. Mann, M. K	2012/05/19 343FraT01	ATGTCTCAA1
56.6177	-3.8837	D. Mann, M. K	2012/05/19 344UlnT01	ATGTCTCAA1
55.895467	-3.308024	D. Mann, M. K	2012/05/19 358UlnW01	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 378FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 379FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 380FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 381FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 382FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 383FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 385FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 387FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 388FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 404FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 409FraB10	ATGTCTCAA1
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56.206051	-4.790649	D. Mann, M. K	2012/05/19 412FraB10	ATGTCTCAA1
56.206051	-4.790649	D. Mann, M. K	2012/05/19 414FraB10	ATGTCTCAA1
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55.964797	-3.20623	D. Mann, M. K	2012/05/19 467FraR03	ATGTCTCAA1
55.964797	-3.20623	D. Mann, M. K	2012/05/19 488FraR03	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 511FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/06/20 513FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/06/20 514FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 515FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 516FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 517FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 551FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 555FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/06/20 575FraK01	ATGTCTCAA1
54.665038	-1.8976458	D. Mann, M. K	2012/05/19 591FraK01	ATGTCTCAA1
56.70746	-3.750611	D. Mann, M. K	2012/05/19 613FraP11	ATGTCTCAA1
56.70746	-3.750611	D. Mann, M. K	2012/06/20 621FraP11	ATGTCTCAA1
56.70746	-3.750611	D. Mann, M. K	2012/05/19 622FraP11	ATGTCTCAA1
56.70746	-3.750611	D. Mann, M. K	2012/06/20 625FraP11	ATGTCTCAA1
51.052806	-1.1697483	D. Mann, M. K	2012/05/19 643FraK06	ATGTCTCAA1
51.194316	-1.4798145	D. Mann, M. K	2012/09/19 653FraK08	ATGTCTCAA1

51.194316	-1.4798145	D. Mann, M. K	2012/05/19	657UlnK08	ATGTCTCAA1
51.215536	-1.4795755	D. Mann, M. K	2012/05/19	662FraK09	ATGTCTCAA1
51.215536	-1.4795755	D. Mann, M. K	2012/05/19	665SynK09	ATGTCTCAA1
51.215536	-1.4795755	D. Mann, M. K	2012/05/19	668SynK09	ATGTCTCAA1
51.420638	-1.3496305	D. Mann, M. K	2012/05/19	698UlnK10	ATGTCTCAA1
51.424326	-1.6739029	D. Mann, M. K	2012/05/19	702UlnK11	ATGTCTCAA1
51.424326	-1.6739029	D. Mann, M. K	2012/05/19	706SynK11	ATGTCTCAA1
51.193109	-2.1752506	D. Mann, M. K	2012/05/19	720UlnK13	ATGTCTCAA1
55.369836	-3.12179	D. Mann, M. K	2012/05/19	726FraB12	ATGTCTCAA1
55.369836	-3.12179	D. Mann, M. K	2012/09/23	732FraB12	ATGTCTCAA1
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55.369836	-3.12179	D. Mann, M. K	2012/05/19	741FraB13	ATGTCTCAA1
55.369836	-3.12179	D. Mann, M. K	2012/05/19	743FraB14	ATGTCTCAA1
55.369836	-3.12179	D. Mann, M. K	2012/05/19	746FraB14	ATGTCTCAA1
55.5185	-2.0174	D. Mann, M. K	2012/05/19	791FraN01	ATGTCTCAA1
55.5185	-2.0174	D. Mann, M. K	2012/05/19	819FraN02	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	820SynN05	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	823SynN05	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	836UlnN05	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	842FraN04	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	844FraN04	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	845FraN04	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	852FraN04	ATGTCTCAA1
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55.51515	-2.044313	D. Mann, M. K	2012/05/19	855FraN04	ATGTCTCAA1
55.51515	-2.044313	D. Mann, M. K	2012/05/19	856FraN04	ATGTCTCAA1
55.964797	-3.20623	D. Mann, M. K	2012/05/19	881UlnR05	ATGTCTCAA1
55.964797	-3.20623	D. Mann, M. K	2012/05/19	907FraR05	ATGTCTCAA1
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55.134333	8.931167	Medlin,L.K. - [2003-07-28	AM713181	AAATGCAAC
55.134333	8.931167	Medlin,L.K. - [2003-07-28	HQ828187	AAATGCAAC
53.55	10.91933	Medlin,L.K. - [2002-07-29	AM710473	AAATGCATC
54.11	10.26	Ruck,E.C. and	1998--	HQ912499	AAAGTGACC
		Ruck,E.C. and	1998--	HQ912454	ATGTCTCAA1
45.896893918	6.1372663652	INRA	1995--	TCC134-Rbcl-1	ATCAAAGTG
45.7296783	5.8695787999	INRA	2008/11/04	TCC301-Rbcl-1	CAGAACGGA
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45.7296783	5.8695787999	INRA	2008/11/04	TCC304-Rbcl-1	AGCTGGTGA
45.7296783	5.8695787999	INRA	2008/11/05	TCC306-Rbcl-1	AAGTGTCGG
46.451413755	6.5732835615	INRA	2009/10/07	TCC365-Rbcl-1	AAGTGACCG
46.3683253216	4538072431	Kermarrec	2009/10/07	TCC367-Rbcl-1	TCTGGTGTA,
-20.95610766	55.298626708	Gilles Gassiole	2010/11/15	TCC520-Rbcl-1	AAAGTGACC
-20.95610766	55.298626708	Gilles Gassiole	2010/11/15	TCC522-Rbcl-1	TCAAAGTGA
-21.026512	55.558742	Gilles Gassiole	2010/11/15	TCC541-Rbcl-1	AGTTCATAC
-21.28301309	55.612390312	Gilles Gassiole	2010/11/15	TCC547-Rbcl-1	TTCATACGCT
-21.28301309	55.612390312	Gilles Gassiole	2010/11/15	TCC553-Rbcl-1	TTCATACGCT
-21.28349295	55.613076957	Gilles Gassiole	2010/11/15	TCC558-Rbcl-1	AGTGACCGT
-20.961426	55.578578	Gilles Gassiole	2010/11/15	TCC559-Rbcl-1	TTCATACGCT
-20.961426	55.578578	Gilles Gassiole	2010/11/15	TCC562-Rbcl-1	TTCATACGCT

48.433333	6.283333	Frédéric Rime	2010/11/15	TCC584-Rbcl-1	AAGTGACCG
47.434958653	-1.494850158	Lenaig Kermar	2010/11/15	TCC589-Rbcl-1	TGACCGTTCC
45.591261735	4.7786832963	Lenaig Kermar	2010/11/15	TCC626-Rbcl-1	GTGTCCGTT/
50.045948056	5.9325362169	Lenaig Kermar	2010/11/15	TCC633-Rbcl-1	CATACGCTG
50.045948056	5.9325362169	Lenaig Kermar	2010/11/15	TCC634-Rbcl-1	CAAGTGTCC
50.045948056	5.9325362169	Lenaig Kermar	2010/11/15	TCC635-Rbcl-1	TTCATACGCT
49.658333	6.137222	Lenaig Kermar	2010/11/15	TCC654-Rbcl-1	CGTTTCAAT
49.816223949	6.0945068359	Lenaig Kermar	2010/11/15	TCC656-Rbcl-1	TGTCCGTTAC
49.932348145	6.2099189682	Lenaig Kermar	2010/11/15	TCC662-Rbcl-1	TCAAGTGAC
49.906485549	5.9591320952	Lenaig Kermar	2010/11/15	TCC666-Rbcl-1	AAAGTGACC
45.419317871	11.018913444	Luc Ector	2010/11/15	TCC669-Rbcl-1	TCAAGTGAC
46.135950021	11.112547126	Luc Ector	2010/11/15	TCC670-Rbcl-1	CCGTTACGA
46.184759184	11.327996650	Luc Ector	2010/11/15	TCC671-Rbcl-1	AAAGTGACC
46.184759184	11.327996650	Luc Ector	2010/11/15	TCC673-Rbcl-1	GACCGTTAC
46.184759184	11.327996650	Luc Ector	2010/11/15	TCC677-Rbcl-1	CAAGTGACC
46.166667	11.3	Luc Ector	2010/11/15	TCC681-Rbcl-1	AAGTGACCG
46.166667	11.3	Luc Ector	2010/11/15	TCC682-Rbcl-1	AAGTGACCG
45.133333	7.15	Luc Ector	2010/11/15	TCC686-Rbcl-1	AAGTGACCG
46.050714824	11.110143867	Luc Ector	2010/11/15	TCC695-Rbcl-1	TAGCATTAT
42.884530468	-4.789637931	Luc Ector	2010/11/15	TCC699-Rbcl-1	ATTAGCATT/
42.417491029	-4.369756904	Luc Ector	2010/11/15	TCC705-Rbcl-1	AATCTGGTG
42.970684050	-4.924550091	Luc Ector	2010/11/15	TCC716-Rbcl-1	GTGTCCGTT/
37.398363538	-8.735049613	Luc Ector	2010/11/15	TCC722-Rbcl-1	AAGTGACCG
37.385106187	-8.649999999	Luc Ector	2010/11/15	TCC728-Rbcl-1	AAAAGTGAC
37.385106187	-8.649999999	Luc Ector	2010/11/15	TCC729-Rbcl-1	AAGTGACCG
46.342577215	6.3792131269	F. Rimet	2010/12/10	TCC743-Rbcl-1	AAAGTGACC
46.342577215	6.3792131269	F. Rimet	2010/12/10	TCC747-Rbcl-1	AATCTGGTG
46.372826226	6.3405427778	F. Rimet	2010/12/10	TCC752-Rbcl-1	GTGACCGTT/
47.094624468	5.8737199628	INRA	2004/05/06	TCC7a-Rbcl-1	TACTGGGAT
46.36718	6.453642	Sylvain Guyot,	2013/01/24	TCC829-Rbcl-1	CCCTTACGCT
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC862-Rbcl-1	CTTAACAGC
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC863-Rbcl-1	CATCGCTTA
59.707064	18.603658	Maria Kahlert,	2013/06/25	TCC865-Rbcl-1	AATCCCTTAC
59.707064	18.603658	Maria Kahlert,	2013/06/25	TCC866-Rbcl-1	AATCTGGTG
59.757285	18.7206	Maria Kahlert,	2013/06/25	TCC867-Rbcl-1	AGCTAACTT/
59.757285	18.7206	Maria Kahlert,	2013/06/25	TCC868-Rbcl-1	ATTATTCCGT
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC869-Rbcl-1	GGACAGATT
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC870-Rbcl-1	ACTGTTGTA
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC871-Rbcl-1	AACAGCTTG
59.707064	18.603658	Maria Kahlert,	2013/06/25	TCC873-Rbcl-1	TATGGACAG
59.707064	18.603658	Maria Kahlert,	2013/06/25	TCC874-Rbcl-1	GAAGCAGCA
59.75728	18.72059	Maria Kahlert,	2013/06/25	TCC877-Rbcl-1	TACTTTGCTT
59.757282	18.720598	Maria Kahlert,	2013/06/25	TCC878-Rbcl-1	ACCGAGTAG
59.707064	18.603658	Maria Kahlert,	2013/07/02	TCC882-Rbcl-1	TGCAACTGA
59.757285	18.7206	Maria Kahlert,	2013/07/02	TCC887-Rbcl-1	GCTTTCATCC

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AM713181 short *rbcl* sequence

HQ828187 short rbcL sequence

AM710473 short rbcL sequence

HQ912499 short rbcL sequence

HQ912454

KT072899 short rbcL sequence

KF959640

0
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KT072900

KT072903

0

KC736614

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KC736594 short rbcL sequence

00

0 short rbcL sequence

0

	Length [μm]	Width [μm]	Striae 10μm ⁻¹	spines	colonies	reference
<i>vaucheriae</i> (Kütz.) J.B.Petersen	14.1-50.4	3.8-5.1	11-14	several conic small spines located along the valve face/mantle junction. Irregularly distributed	never more than two loosely attached cells could be found as also observed by Petersen (1938)	Wetzel and Ector (2015) ^t
<i>vaucheriae</i>	no info	no info	9-14	no info	ribbon like, attached whole way	Tuji and Williams (2006a) ^t
<i>vaucheriae</i>	no info	no info	no info	spines either do not exist or are very small	do not link with sibling cells	Tuji and Williams (2013) ^t
<i>vaucheriae</i>	<10-50	4-5	9-14	no info	no info	Hofmann et al. (2011)
<i>intermedia</i> (synonym to <i>vaucheriae</i>)	no info	no info	no info	SEM shows none, or very small spines	SEM shows single cells only	Tuji and Williams (2013) ^t Note: synonymized <i>F. intermedia</i> with <i>F. vaucheriae</i>
<i>intermedia</i> (Grunow) Grunow in van Heurck (1881)	16-30	3-4	11-14	no info	no info	Delgado et al. (2015), analyzing figures of Tuji and Williams (2013) ^t Note: refer to Tuji and Williams (2013), but still list <i>F. intermedia</i> as valid name
<i>neointermedia</i> Tuji & Williams	25-35 ^t	3.5-4.5 ^t	8-10 ^t	spathulate linking	long colonies	Tuji and Williams (2013) ^t
<i>neointermedia</i>	25-35 ^{LM} 23-29 ^{SEM}	3.5-4.5 ^{LM} 3-4 ^{SEM}	8-10 ^{LM} 10-12 ^{SEM}	no info	no info	Delgado et al. (2015), re-analyzing figures in Tuji and Williams (2013) ^t
<i>neointermedia</i>	22.5-45	3.4-4.2	11-14	spathulate linking	no info	Delgado et al. (2015) ^t , analyzed type material
<i>capucina</i> Desmazières s.str.	24-65 ^f	3-5 ^f	13-17 ^f	linking spines	ribbon like; (Tuji and Williams (2006b) write it is attached)	Delgado et al. (2015) ^t re-analyzing figures in Tuji and Williams (2006b) (2 RP per valve)
<i>capucina</i> Desmazières s.str.	28-47	3.3-4.2	14-17	linking spines	no info	Delgado et al. (2015) ^t (2 RP per valve)
" <i>capucina</i> " 1 (aff.)	33.2-35.2	3.2-3.4	15	"lacking" marginal spines	no info	Tuji and Williams (2006b) ^{t, f}
" <i>capucina</i> " 2 (aff.)	18-20	3.4-3.6	15	no info in text; no spines visible on SEM (Tuji and Williams 2006b). (The info by Delgado et al. (2015) ^t of "linking spines" is probably a misinterpretation of Tuji & Williams (2006b)'s text).	no info	Tuji and Williams (2006b) ^{t, f}

<i>capucina</i>	20-75	3.5-4.5	12-17	no info	ribbon-bands OR stellate	Hofmann et al. (2011), discussing Krammer and Lange-Bertalot (1991), probably including all 3 types in typeslide
<i>microvaucheriae</i> C.E.Wetzel & Ector	5.7-23.4	2.5-3.8	15-16	small, conic. Usually absent	no colonies	Wetzel and Ector (2015) ^f
<i>pectinalis</i> (O.F.Müll.) Lyngb.	12.0-36.2	2.8-4.7	15-18	totally absent	no colonies	Wetzel and Ector (2015)
<i>pectinalis</i>	28-37	3,5-4	14-15	a few, very small, spines, visible in SEM	No information, but SEM shows a pair of cells	Tuji and Williams (2006b), Tuji and Williams (2008b) ^f
<i>rinoi</i> Almeida & C.Delgado	8.8-24.1	4.2-5.6	14-16	totally absent	no colonies	Delgado et al. (2016) ^f
<i>uliginosa</i> Kulikovskiy, Lange-Bert., Witkowski & Dorofeyuk	14.1-28.8	3.5-4.5	15-16 ^f 16-17 (table)	wide, spathulated and usually located on the striae at the valve face/mantle junction	colonies were not observed and the species seems to be mainly found as solitary cells	Wetzel and Ector (2015) ^f
<i>recapitellata</i> Lange- Bert. & Metzeltin	20-39 ^f 21-37 ^f	3-4 ^f 3-4 ^f	17-19 ^f 18-20 ^f	no spines ^f	no info	Tuji and Williams (2008b), analysis of figures by Delgado et al. (2015)
<i>recapitellata</i>	21.4-30.4	2.8-4.2	17-19	no info	no info	Delgado et al. (2015), analyzed type material
<i>recapitellata</i>	11-38	3-5	14-18	no info		Hofmann et al. (2011)
<i>recapitellata</i>	15-27	4.1-5.6	16-20	no spines	no info	Bishop and Spaulding (2014)* (*"Note that the taxon illustrated here conforms more to the taxon illustrated in Metzeltin et al. (2009) than in the original description of <i>Synedra capitellata</i> Grunow in Van Heurck 1881")
<i>perminuta</i> (Grunow) Lange-Bert.	9-24	3-3.5	18-19	no material for SEM left, thus no info	no info	Tuji and Williams (2008b) ^f
<i>perminuta</i>	8-25 ^f	3-4 ^f	17-19 ^f	no info	no info	Delgado et al. (2015), re-analysed Tuji and Williams (2008b) ^f
<i>perminuta</i>	7-40	3-4	17-21	no info	no info	Hofmann et al. (2011)
<i>mesolepta</i> Rabenhorst	23.8-55.9 ^f	3.9-5 ^f	13-15 ^f	spathular linking	ribbon closed to ends	Tuji and Williams (2008a) ^f (note: RP on mantle/valve junction)
<i>mesolepta</i>	20-60	3.5-4.5	15-18	no info	no info	Hofmann et al. (2011) ^f
<i>tenuistriata</i> Østrup	50.4-74.8 ^f	3.9-4 ^f	15-16 ^f	spathular linking	ribbon closed to ends	Tuji and Williams (2008a) ^f (note: RP on sternum)

<i>subconstricta</i> Østrup	39.4-47.4 ^f	3.7 ^f	15-16 ^f	spathular linking	ribbon closed to ends	Tuji and Williams (2008a) ^f (notes: RP on mantle/valve junction, but different valve shape than <i>F. mesolepta</i>)
<i>rhabdosoma</i> Ehrenberg	~27	~3-4	15	spathular, nicely intertwined between cells	long ribbon bands	Tuji (2004) (considers <i>F. bidens</i> being a synonym to <i>F. rhabdosoma</i>)
<i>"bidens"</i> Heiberg	10-50	(2)3-4	(11?)15-18	no info	long ribbon bands	Krammer and Lange-Bertalot (1991), (complicated taxonomy and synonymy, including also <i>F. socia</i> , <i>F. familiaris</i> , <i>F. parva</i> , <i>F. rhabdosoma</i> , see also Lange-Bertalot (1980))
<i>radians</i> (Kütz.) Lange-Bert.	35-55	3.5-4.5	9-11	no SEM pictures	radial colonies ("stellate")	Krammer and Lange-Bertalot (1991), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>austriaca</i> (Grunow) Lange-Bert.	20-60	3-4	12-15	no SEM pictures	ribbon-bands OR stellate	Krammer and Lange-Bertalot (1991), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>amphicephaloides</i> Lange-Bert.	40-75	2-3	10-14	no SEM pictures	ribbon-bands OR stellate	Krammer and Lange-Bertalot (1991), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>henryi</i> Lange-Bert.	35-60	3-4	11-13p	no SEM pictures?	ribbon-bands OR stellate OR singular cells	Krammer and Lange-Bertalot (1991), Lange-Bertalot and Genkal (1999), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>acidoclinata</i> Lange-Bert. & G. Hofmann	35-60	3-4	11-13p	no SEM pictures?	ribbon-bands OR stellate OR singular cells	Krammer and Lange-Bertalot (1991), Lange-Bertalot (1993), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>famelica</i> (Kütz.) Lange-Bert.	10-70	2.5-4	11-16p	no SEM pictures?	ribbon-bands OR stellate OR singular cells	Krammer and Lange-Bertalot (1991), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>pararumpens</i> Lange-Bert., G. Hofmann & Werum	25-50	2.5-3	16-18	linking spines, thorn-formed with cylindric base and flatted anchors, which are tightly connected at neighboring frustules; at the valve ends spines acute thorn-formed, not connecting	chain formed connected in the middle (double-comb), similar to <i>F. crotonensis</i>	Krammer and Lange-Bertalot (1991), Krammer and Lange-Bertalot (2004), Hofmann et al. (2011)
<i>parva</i> Tuji & D. M. Williams	30-40 ^f	~3 ^f	~20 ^f	no info	no info	Tuji and Williams (2008c)

<i>socia</i> (Wallace) Lange-Bert.	16-72	3.5-4	17	no material for SEM left, thus no info	no info	Lange-Bertalot (1980), Tuji and Williams (2008b)
<i>socia</i>	14.1-42.9	3.4-4.1	16.4-18.5	no info	frustules are joined in colonies, attached to a benthic substrate by a mucilage pad (pictures: stellate)	LaLiberte and Vaccarino (2015)
<i>rumpens</i> (Kütz.) G. W. F. Carlson	25-63	3-4	18-20	irregular, located on the costae, at mantle-face junction, often deformed and rectangular at central area, very small and triangular at valve poles	ribbon like, adhering by valve faces, sometimes separated from each other at poles (tychoplanktonic to attached)	Tuji and Williams (2006a) ^t
<i>rumpens</i>	20-65	3.5-4.5	18-20	no info	ribbon band or stellate adnate	Hofmann et al. (2011)
<i>gracilis</i> Østrup	36	2-3/3.6	20p	absent	no info	Tuji (2007)
<i>gracilis</i>	no info	no info	18-20p	absent	no info	Schmidt et al. (2004)
<i>gracilis</i>	30-50	2-3	20-22*	absent	no info	Lange-Bertalot and Ulrich (2014) (* "alternating and opposing in the same valve")
<i>gracilis</i>	10-60	2-3	~20	no info	ribbon-bands OR stellate	Hofmann et al. (2011)
<i>aquaplus</i> Lange-Bert. & S. Ulrich*	30-50 22-45 (table)	1.5-2.5	22-24p	absent	solitary	Lange-Bertalot and Ulrich (2014) (*this taxon is one of the 2 taxa on the <i>Synedra nana</i> Meister 1912 typeslide. Was pooled by Krammer and Lange-Bertalot (1991) into <i>F. nanana</i> Lange-Bert.)
<i>nanana</i> sensu Lange-Bertalot (1991)	40-90	1.5-2	22-30	no info	no info	Krammer and Lange-Bertalot (1991)
<i>nanana</i> sensu Lange-Bertalot (1991)	40-90	1.5-2	22-30*	no info	no info	Hofmann et al. (2011) (* „striae coaxial, not alternating“)
<i>nanana</i> sensu Lange-Bertalot (1991)	no info	1.5-2	no info	absent	no info	Schmidt et al. (2004)
<i>tenera</i> var. <i>nanana</i> (Lange-Bertalot)	29-85 (table)	2-2.3 (table)	18.5–20	present marginal and apical	no colonies	Lange-Bertalot and Ulrich (2014) (*this taxon is one of the 2 taxa on the <i>Synedra nana</i> Meister 1912 typeslide. Was pooled by Krammer and Lange-Bertalot (1991) into <i>F. nanana</i> Lange-Bert.)
<i>saxoplanctonica</i> Lange-Bert. & Ulrich	40–170	1.5–2.5	23–28 pa	absent	single individuals, not in ribbon-like or stellate colonies	Lange-Bertalot and Ulrich (2014)

<i>sepes</i> Ehrenberg				small rectangular spines	yes, ribbon-like; Ehrenberg picture shows a kind of long ribbon band, could also be a double-comb	Tuji (2004)
<i>sepes</i>	47.0–69.0	1.5–2.0	24–25 pa	small, pyramidal	nd.	Almeida et al. (2016), re-analyzed Tuji (2004)
<i>tenuissima</i> Lange-Bert. & Ulrich	40–150	1–3	16–20.5	marginal spines may be reduced	may form loose fewcelled aggregates	Lange-Bertalot and Ulrich (2014)
<i>tenuissima</i>	70–145	1.6–2.8	16.0–20.5	small, reduced or absent	single, or short ribbon-like	Almeida et al. (2016) re-analyzed Lange-Bertalot and Ulrich (2014)
<i>tenera</i> (W. Smith) Lange-Bert.	68.1–114.4	1.9–2.1	18–20	pyramidal spines	at most loose aggregates	Almeida et al. (2016)
<i>tenera</i> var. <i>tenera</i>	60–120	1.8–2.5	18–20	marginal and apical	may form loose fewcelled aggregates	Lange-Bertalot and Ulrich (2014)
<i>tenera</i>	30–>100	2–3	17–20	no info	no info	Hofmann et al. (2011)
<i>tenera</i>	30–100	2–3	17–20	no info	never forms colonies, always solitary	Druart et al. (2007)
<i>tenera</i> var. <i>lemanensis</i> Druart, Lavigne & Robert	70–80	2–3.5	18–20	no info	stellate	Druart et al. (2007)
<i>tenera</i> var. <i>lemanensis</i>	70–90 (table)	2–3.5 ^t	18–20 ^t	marginal and apical	stellate	Lange-Bertalot and Ulrich (2014)
<i>nanoides</i> Lange-Bert.	40–90	1.8–2.4	22.5–24a	no info	no info	Lange-Bertalot (1996)
<i>nanoides</i>			>20	absent	occurrence of cell pairs	Schmidt et al. (2004)
<i>spectra</i> P.D.Almeida, E.Morales & C.E.Wetzel	40.5–73	1.5–2.5	24–25	absent	solitary	Almeida et al. (2016) (2 RP per valve)
<i>neotropica</i> P.D.Almeida, E.Morales & C.E.Wetzel	52–72	1.7–2	28–32	pyramidal, at junction valve/mantle	not observed	Almeida et al. (2016)
<i>longifusiformis</i> ssp. <i>longifusiformis</i> (Hains & Sebring) Siver et al.	50–175	2–4	26–34 pa	reduced marginal spines, few apical spines of different size	no info	Lange-Bertalot and Ulrich (2014)
<i>longifusiformis</i> ssp. <i>eurofusiformis</i> Lange-Bert. & S.Ulrich	70–120	3–4	29–31 pa	reduced marginal spines, few apical spines of different size	very rarely, bunch-like	Lange-Bertalot and Ulrich (2014)
<i>grunowii</i> Lange-Bert. & S.Ulrich	90–380	3–4	12–15 pa	no, at most small blunt apical dents	solitary	Lange-Bertalot and Ulrich (2014)

<i>schoeteri</i> (Meister) Lange-Bert. & S.Ulrich	300-450	2.5–4.5	12–15	none marginal, but 2 short apical	solitary	Lange-Bertalot and Ulrich (2014)
<i>paludosa</i> (Meister) Lange-Bert. & S.Ulrich	73–110	3.2-3.9	14-15	no info	no info	Lange-Bertalot and Ulrich (2014)
<i>delicatissima</i> (W.Smith) Lange-Bert.	30-100	2.5-3	14-16	no info	no info	Krammer and Lange-Bertalot (1991), Hofmann et al. (2011)
<i>perdelicatissima</i> Lange- Bert. & Van de Vijver	36-95	2-2.6	14-16	absent	Solitary (benthic)	Lange-Bertalot and Ulrich (2014), <i>F. delicatissima</i> (W. Smith) Lange-Bertalot sensu Krammer & Lange-Bertalot 1991, p. 129, fig. 115: 13
<i>crotonensis</i> ssp. <i>crotonensis</i> Kitton	40-170	2-4(5), mostly 2.5-3.5	15–18	spatula shaped linking spines, marginal spines short & acute (but no single valves with short, acute spines)	ribbon-like colonies, rarely single cells, cells connected only in the proximal inflated part	Lange-Bertalot and Ulrich (2014)
<i>crotonensis</i> ssp. <i>lacus- vulcani</i> Lange-Bert. & S.Ulrich	55-120	2-4 (long forms: 2-3, short forms 3- 4)	14-17	spatula shaped linking spines, marginal spines short & acute (plus many single valves with short acute marginal spines)	cells mainly single or in few-celled aggregates	Lange-Bertalot and Ulrich (2014)
<i>crotonensis</i> "rod-form": <i>F. crotonensis</i> var. <i>crotonensis</i> Kitton	34-100, rod- formed ends in girdle view	No info	14-15 center, 17-18 in apices	Interlocking spines are narrow at the base and broaden towards their spatulate tips, which interdigitate with those of the sibling valve. Outside the linking zone, the spines change in form, taper distally, and tend to slant toward the cell apex. Valves at the ends of intact filaments lack a linking zone and may be regarded as separation valves. On these valves all of the spines are tapered, simple, and small.	long raft-like chains, with wide mucilage envelope (up to 20 µm)	Crawford et al. (1985) (did not formally describe the 2 forms), difference to flared-form in parasite suceptibility
<i>crotonensis</i> "flared form": <i>F. crotonensis</i> var. <i>prolongata</i> Grunow ex van Heurck.	58-92 flared- formed ends in girdle view	No info	14-15 center, 17-18 in apices	same as before	long raft-like chains, with very narrow mucilage envelope (1-6 µm)	Crawford et al. (1985) (did not formally describe the 2 forms), difference to flared-form in parasite suceptibility

References

Almeida, P. D., Morales, E. A., Wetzel, C. E., Ector, L. & Bicudo, D. C. 2016. Two new diatoms in the genus *Fragilaria* Lyngbye (Fragilariophyceae) from tropical reservoirs in Brazil and comparison with type material of *F. tenera*. *Phytotaxa* 246:163–83.

Bishop, I. & Spaulding, S. 2014. *Fragilaria recapitellata*. Available at: https://diatoms.org/species/fragilaria_capucina_var._capitellata (last accessed July 30 2018).

Crawford, R. M., Canter, H. M. & Jaworski, G. H. M. 1985. A study of 2 morphological variants of the diatom *fragilaria-crotonensis* kitton using electron-microscopy. *Annals of Botany* 55:473-85.

Delgado, C., Novais, M. H., Blanco, S. & Almeida, S. F. P. 2015. Examination and comparison of *Fragilaria candidagilae* sp. nov. with type material of *Fragilaria recapitellata*, *F. capucina*, *F. perminuta*, *F. intermedia* and *F. neointermedia* (Bacillariophyta, Fragilariaceae). *Phytotaxa* 231:001-18.

Delgado, C., Novais, M. H., Blanco, S. & Almeida, S. F. P. 2016. *Fragilaria rinoi* sp nov (Fragilariales, Fragilariophyceae) from periphytic river samples in Central Portugal. *Eur. J. Taxon.* 248:1-16.

Druart, J. C., Lavigne, S. & Robert, M. 2007. *Fragilaria tenera* var. *lemanensis*, a new variety from the lake of Geneva (France, Switzerland). *Cryptogamie Algologie* 28:283-87.

Hofmann, G., Werum, M. & Lange - Bertalot, H. 2011. *Diatomeen im Süßwasser-Benthos von Mitteleuropa. Bestimmungsflora Kieselalgen für die ökologische Praxis. Über 700 der häufigsten Arten und ihre Ökologie.* A.R.G.Gantner Verlag K.G., Rugell, 908.

Krammer, K. & Lange-Bertalot, H. 1991. *Bacillariophyceae. 3. Teil: Centrales, Fragilariaceae, Eunotiaceae. Süßwasserflora von Mitteleuropa.* Gustav Fischer Verlag, Stuttgart New York, 576.

Krammer, K. & Lange-Bertalot, H. 2004. *Bacillariophyceae. 3. Teil: Centrales, Fragilariaceae, Eunotiaceae. Rev. ed. 2004. With new supplement (= p. 580-599). Reprint 2008.* Gustav Fischer Verlag, Stuttgart New York, 599.

LaLiberte, G. & Vaccarino, M. 2015. *Fragilaria socia*. In *Diatoms of North America.* . Available at: https://diatoms.org/species/fragilaria_socia (last accessed 28 August 2018).

Lange-Bertalot, H. 1980. Zur systematischen Bewertung der bandförmigen Kolonien bei *Navicula* und *Fragilaria*. *Nova Hedwigia* 33:723.

Lange-Bertalot, H. 1993. *85 Neue Taxa und über 100 weitere neu definierte Taxa ergänzend zur Süßwasserflora von Mitteleuropa, vol. 2/1-4.* J. Cramer, Berlin, 1-428.

Lange-Bertalot, H. 1996. *Annotated Diatom Micrographs Vol. 2. Indicators of Oligotrophy, by Lange-Bertalot, H. & Metzeltin.* D. Koeltz Scientific Books,

Lange-Bertalot, H. & Genkal, S. I. 1999. *Diatoms of Siberia.* Koeltz Scientific Books, Königstein,

Lange-Bertalot, H. & Ulrich, S. 2014. Contributions to the taxonomy of needle-shaped *Fragilaria* and *Ulnaria* species. *Lauterbornia* 78:1-73.

Petersen, J. B. 1938. *Fragilaria intermedia-Synedra vaucheriae.* *Botaniska Notiser* 1938:164.

Schmidt, R., Kamenik, C., Lange-Bertalot, H. & Klee, R. 2004. *Fragilaria* and *Staurosira* (Bacillariophyceae) from sediment surfaces of 40 lakes in the Austrian Alps in relation to environmental variables, and their potential for palaeoclimatology. *2004* 63:19.

- Tuji, A. 2004. Type and examination of the ribbon-forming *Fragilaria capucina* complex described by Christian Gottfried Ehrenberg. . In: Poulin, M. [Ed.] *Proceeding of the Seventeenth International Diatom Symposim*. Biopress Limited, Bristol, England. pp. 411-422., pp. 411-22.
- Tuji, A. 2007. Type examination of *Fragilaria gracilis* Østrup (Bacillariophyceae). *Bulletin of the national museum of nature and science. Series B, Botany* 33:9-12.
- Tuji, A. & Williams, D. M. 2006a. Examination of the type material of *Synedra rumpens* = *Fragilaria rumpens*, Bacillariophyceae. *Phycological Research* 54:99-103.
- Tuji, A. & Williams, D. M. 2006b. Typification of *Conferva pectinalis* O.F. Müll. (Bacillariophyceae) and the identity of the type of an alleged synonym. *Fragilaria capucina* Desm. *Taxon* 55:193.
- Tuji, A. & Williams, D. M. 2008a. Examination of type material of *Fragilaria mesolepta* Rabenhorst and two similar, but distinct, taxa. *Diatom Research* 23:503-10.
- Tuji, A. & Williams, D. M. 2008b. Examination of types in the *Fragilaria pectinalis-capitellata* species complex. In: Likhoshway, Y. [Ed.] *Proceedings of the Nineteenth International Diatom Symposium 2006, Listvyanka, Russia*. Biopress Limited, Bristol, Listvyanka, Russia, pp. 125-39.
- Tuji, A. & Williams, D. M. 2008c. Typification and type examination of *Synedra familiaris* Kütz. and related taxa. *Diatom. The Japanese Journal of Diatomology* 24:25-29.
- Tuji, A. & Williams, D. M. 2013. Examination of types in the *Fragilaria vaucheriae-intermedia* species complex. *Bulletin of the national museum of nature and science Series B, Botany* 39:1.
- Wetzel, C. E. & Ector, L. 2015. Taxonomy and Ecology of *Fragilaria microvaucheriae* sp. nov. and Comparison with the Type Materials of *F. uliginosa* and *F. vaucheriae*. *Cryptogamie, Algologie* 36:271-89.

Table S3. Towards more harmonized taxa names: Suggestions on how to separate Fragilaria species by morphological characters in LM (based on published analyses of type material). Fet style: typical character to use for separation in LM.

FRAGILARIA	length [µm]	width [µm]	striae 10µm ⁻¹	colonies*	characters	reference	note
"medium-sized"							
<i>F. neointermedia</i> Tuji & Williams	23-35	3-4.5	8-12	yes	few striae	1	
<i>F. vaucheriae</i> (Kütz.) J.B.Petersen	14.1-50.4	3.8-5.1	11-14	no	few striae	2	
<i>F. rinoi</i> Almeida & C.Delgado	12-36	4.2-5.6	14-16	no	SEM: no spines	1	" <i>F.pectinalis</i> s.lat. without colonies" - complex not separable in LM
<i>F. uliginosa</i> Kulikovskiy, Lange-Bert., Witkowski & Dorofeyuk	14.1-28.8	3.5-4.4	15-17	no	SEM: large linking spines	2	" <i>F.pectinalis</i> s.lat. without colonies" - complex not separable in LM
<i>F. microvaucheriae</i> C.E.Wetzel & Ector	5.7-23.4	2.5-3.8	15-16	no	< 4µm wide; L:W 2-6, SEM: tiny or no spines	2	" <i>F.pectinalis</i> s.lat. without colonies" - complex not separable in LM
<i>F. pectinalis</i> (O.F.Müll.) Lyngb.	12-36	2.8-4.7	15-18	no	L:W 6-8, SEM: tiny spines	2, 4	" <i>F.pectinalis</i> s.lat. without colonies" - complex not separable in LM
<i>F. heatherae</i> sp. nov. (FCAP1)	9-38	3.3-3.7	16(18)	± no	< 4µm wide; L:W up to 11, SEM: no spines	13	" <i>F.pectinalis</i> s.lat. without colonies" - complex not separable in LM
<i>F. capucina</i> s.str. Desm.	28-47	3.3-4.2	14-17	yes	SEM: large linking spines	3, 1	" <i>F.capucina</i> s.lat. with colonies" - complex not separable in LM
<i>f. joachimii</i> sp. nov. (FCAP2)	5-35	3.3-4.6	14-16	yes	L:W ratio 5.3 to 9.4, SEM: tiny spines	13	" <i>F.capucina</i> s.lat. with colonies" - complex not separable in LM
<i>F. agnesiae</i> sp. nov (FVAU)	9-65	4.0-5.4	14-16	yes	SEM: tiny spines	13	" <i>F.capucina</i> s.lat. with colonies" - complex not separable in LM
<i>F. perminuta</i> (Grunow) Lange-Bertalot	8-25	3-4	17-21	no info	< 4µm wide, rhombic valve form, rimmed central unilateral depression/swelling	1, 6, 7	
<i>F. recapitellata</i> Lange-Bertalot&Metzeltin	20-39	2.8-4.2	17-20	no info	capitate ends	1, 4, 5	
"medium-long, thin, with ~20 striae 10 µm ⁻¹ⁿ							
<i>F. gracilis</i> Østrup	30-50	2-3	20-22	no	opposite, parallel striae; valve form ± linear	8, 9	
<i>F. tenera</i>	60-120	1.8-2.5	18-20	no	valves linear-lanceolate with only slightly convex margins, subcapitate	9, 10	
<i>F. rumpens</i> (Kütz.) G. W. F. Carlson	25-63	3-4	18-20	yes	width > 3µm	11	
<i>F. pararumpens</i>	25-50	2.5-3	16-18	yes	central swelling, subcapitate ends	7	
<i>F. saxoplanctonica</i>	40-170	1.5-2.5	23-28	no		7, 9	
<i>F. nanoides</i>	40-90	1.8-2.4	22.5-23	no	subcapitate, getting thinner from middle to ends; central area 'empty'	12	
<i>F.aquaplus</i> Lange-Bert. & S. Ulrich	22-45 ⁹	1.5-2	22-24	no	opposite, parallel very delicate striae; valve form ± needle-formed, rounded ends	6, 7, 9, 12, 13	suggestion to synonymize <i>F.aquaplus</i> with <i>F.gracilis</i> , because not separated by rbcl & presence of intermedia forms former called " <i>F. nanana</i> " sensu Lange-Bert. 1991, part of the mixture of " <i>Synedra</i> nana Meister 1912"
<i>F. nanana</i> sensu Lange-Bertalot (1991)	40-90	1.5-2	22-30	no info	striae not alternating	6, 7, 12	part of the mixture of " <i>Synedra nana</i> Meister 1912", name should not be used further

* the presence of colony-formation needs to be analysed in adequately prepared slides

Literature used:

1 Delgado et al. 2015. Phytotaxa 231:001-18.
2 Wetzel & Ector 2015. Cryptogamie, Algologie 36:271-89.
3 Tuji & Williams 2006. Taxon 55:193.
4 Tuji & Williams 2008. In: Likhoshway, Y. [Ed.] Proceedings of the Nineteenth International Diatom Symposium 2006. Biopress Limited.
5 Tuji & Williams 2013. Bull. Natl. Mus. Nat. Sci., Ser. B 39(1):1-9.
6 Krammer & Lange-Bertalot 1991. Bacillariophyceae. 3. Teil. Süßwasserflora von Mitteleuropa. Gustav Fischer Verlag.
7 Hofmann et al. Diatomeen im Süßwasser-Benthos von Mitteleuropa. A.R.G.Gantner Verlag K.G..
8 Tuji 2007, Bull. Natl. Mus. Nat. Sci. Ser. B 33(1):9-12
9 Lange-Bertalot & Ulrich 2014. Lauterbornia 78:1-73.
10 Almeida et al. 2016. Phytotaxa 246:163–83.
11 Tuji & Williams 2006, Phycological research, 54: 99-103
12 Lange-Bertalot 1996. Annotated Diatom Micrographs Vol. 2. Indicators of Oligotrophy. D. Koeltz Scientific Books.
13 this study

Fig. S1. Phylogeny of the studied strains of *Fragilaria*, rooted with strains of the genus *Ulnaria*, on a 1087 bp long part of the *rbcL* barcode. Bootstrap values above 69% are given for each node. Scale bar: number of substitutions per site. Full tree with geographical information included.

